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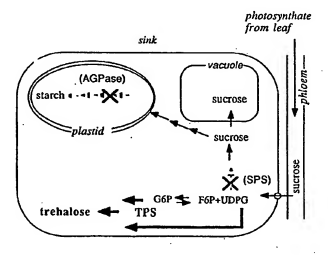
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(54) Title: PRODUCTION OF TREHALOSE IN PLANTS

ENGINEERING OF TREHALOSE-PRODUCTION IN PLANTS



(57) Abstract

The present invention provides for the production of trehalose in a plant host due to the presence in said plant host of a plant expressible gene which comprises in sequence: (a) a transcriptional initiation region that is functional in said plant host; (b) a DNA sequence encoding a trehalose phosphate synthase activity; and optionally, (c) a transcriptional termination sequence that is functional in said plant host.

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PRODUCTION OF TREHALOSE IN PLANTS FIELD OF THE INVENTION

5 This invention relates to the modification of plant carbohydrate metabolism using recombinant DNA techniques, recombinant DNA for use therein, as well as plants and parts of plants having a modified genetic constitution. Said plants may be used to extract specific carbohydrate compounds, or alternatively, they may be processed as food, feed, or ingredients thereof, having improved properties due to the presence of said carbohydrate compounds, <u>e.g.</u> during processing.

STATE OF THE ART

Trehalose is a general name given to D-glucosyl D-glucosides which comprise disaccharides based on two α-, α,β- and β,β-linked glucose molecules. Trehalose, and especially α-trehalose 1-(0-a-D-glucopyranosyl)-1'-0-α-D-glucopyranose) is a widespread naturally occurring disaccharide.

The chemical synthesis of trehalose is difficult (protecting groups required) and inefficient. Current natural sources of trehalose are mushrooms and the yeast Saccharomyces <u>cerevisiae</u>, that can accumulate over 10% of dry 25 weight as trehalose. However production is hampered by high trehalase activity causing rapid metabolization of trehalose. Elbein A.D. (1974, Adv. Carbohydrate Chem. and Biochem. 30, 227-256) gives a review of the occurrence and metabolism of the disaccharide trehalose, particularly α, α -trehalose, in 30 living organisms. In plants, the presence of trehalose has been reported in some lower plant species, as well as in a number of higher plant species belonging to the spermatophyta; Echinops persicus, Carex brunescens; Fagus silvaticus. However, these results have never been firmly 35 established by other authors (e.g. Kendall et al., 1990, Phytochemistry 29, No. 8, 2525-2528). For instance, Kendall et al, supra, referring to the occurrence of trehalose in spermatophytes, stated that the presence thereof has only been firmly documented for caraway seed (Carum carvi). A 40 report of the presence of trehalose in sunflower by Cegla et

al., (1977, J. Am. Oil Chem. Soc. 54, 150 et seq.) was

questioned by Kandler et al., (in: The Biochemistry of Plants

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Vol. 3 Carbohydrates: Structure and Function; Preiss, J., ed., p.228. Academic Press) according to Kendall et al, 1990, supra. Reports of trehalose in beech (Fagus sylvaticus) and cabbage could not be verified by other authors (Kendall et al., 1990, supra, and references therein).

In spite of the apparent rarity of trehalose in higher plants, the presence of trehalose degrading activities was reported for a significant number of the investigated plant families. Stable high trehalase activity was found in three wheat lines, jack pine, and <u>Selaginella lepidophylla</u>. Stable, low trehalase activity was found in alfalfa, black Mexican sweet corn and white spruce. Labile, moderate activities were found in two different suspensions of canola, but these could probably not be ascribed to specific trehalase activity.

Barley, brome grass, soybean and black spruce were reported to contain no trehalase activity at all (Kendall, 1990, supra).

In organisms capable of its production trehalose is believed to be biosynthesized as the 6-phosphate, whereas the storage form is the free sugar. It is therefore believed, that organisms that produce and/or store trehalose contain a phosphatase capable of cleaving trehalose 6-phosphate. (Elbein, 1974, supra). Little is known about the presence of specific trehalose phosphate phosphatases in higher plants.

25

SUMMARY OF THE INVENTION

The present invention provides for a method for the production of trehalose in a plant host due to the presence in said plant host of a plant expressible gene which comprises in sequence:

- (a) a transcriptional initiation region that is functional in said plant host,
- (b) a DNA sequence encoding a trehalose phosphate synthase activity, and optionally
- 35 (c) a transcriptional termination sequence that is functional in said plant host.

Another embodiment of the invention comprises the production of trehalose in a plant host due to the presence in said plant host of a plant expressible gene which

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comprises in sequence:

- . (a) a transcriptional initiation region that is functional in said plant host,
- (b) a DNA sequence encoding a trehalose phosphate synthase 5 activity, and optionally
 - (c) a transcriptional termination sequence that is functional in said plant host, and
 - a plant expressible gene comprising in sequence:
- (a) a transcriptional initiation region that is functional insaid plant host,
 - (b) a DNA sequence encoding an RNA sequence which is at least partially complementary to an RNA sequence which encodes a sucrose phosphate synthase enzyme (SPS) naturally occurring in said plant host, and optionally
- 15 (c) a transcriptional termination sequence that is functional in said plant host.

Yet another embodiment of the invention comprises the production of trehalose in a plant host due to the presence in said plant host of a plant expressible gene which

- 20 comprises in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding a trehalose phosphate synthase activity, and optionally
- 25 (c) a transcriptional termination sequence that is functional in said plant host, and
 - a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
- 30 (b) a DNA sequence encoding an RNA sequence which is at least partially complementary to an RNA sequence which encodes an ADP-glucose pyrophosphorylase enzyme naturally occurring in said plant host, and optionally
- (c) a transcriptional termination sequence that is functional in said plant host.

Yet another embodiment of the invention comprises the production of trehalose in a plant host due to the presence in said plant host of a plant expressible gene which comprises in sequence:

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(a) a transcriptional initiation region that is functional in said plant host,

- (b) a DNA sequence encoding a trehalose phosphate synthase activity, and optionally
- 5 (c) a transcriptional termination sequence that is functional in said plant host,
 - and a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
- 10 (b) a DNA sequence encoding an RNA sequence at least partially complementary to an RNA sequence which encodes a sucrose phosphate synthase enzyme naturally occurring in said plant host, and optionally
- (c) a transcriptional termination sequence that is functionalin said plant host,
 - and a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding an RNA sequence at least
- 20 partially complementary to an RNA sequence which encodes an ADP-glucose pyrophosphorylase enzyme naturally occurring in said plant host, and optionally
 - (c) a transcriptional termination sequence that is functional in said plant host.
- The invention also extends to the plant expressible genes used in the process for making trehalose, as well as to the combinations of plant expressible genes, as well as to cloning plasmids, transformation vectors, microorganisms, an individual plant cells harboring plant expressible genes according to the invention.

The invention also provides a recombinant plant DNA genome which contains a plant expressible trehalose phosphate synthase gene that is not naturally present therein. The invention also comprises a recombinant plant DNA genome which comprises a plant expressible trehalose phosphate gene that is not naturally present therein and in addition a plant expressible gene capable of inhibiting biosynthesis of an SPS activity, and/or a plant expressible gene capable of inhibiting biosynthesis of an AGPase activity.

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The invention also provides a method for obtaining a plant capable of producing trehalose comprising the steps of,

- (1) introducing into a recipient plant cell a plant expressible gene comprising in sequence:
- 5 (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding a trehalose phosphate synthase activity,
- (c) a transcriptional termination sequence that is functional in said plant host, and a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding a selectable marker gene that is functional in said plant host, and optionally
 - (c) a transcriptional termination sequence that is functional in said plant host,
- (2) generating a plant from a transformed cell under conditions that allow for selection for the presence of the 20 selectable marker gene.

The invention also comprises plants which produce (increased levels of) trehalose as a result of genetic modification.

The invention further comprises plants having a 25 recombinant DNA genome containing a plant expressible gene according to the invention.

The invention also comprises plants having a recombinant DNA genome containing a plant expressible gene according to the invention and which plants produce trehalose.

The invention also comprises plants having a recombinant DNA genome according to the invention and which exhibit increased drought resistance.

The invention also extends to parts of plants according to the invention such as cells or protoplasts or cultures

35 thereof, flowers, fruits, leaves, pollen, roots (including hairy root cultures), seeds, stalks, tubers (including so-called microtubers) and the like.

The invention also extends to a method of preserving plants or plant parts in the presence of trehalose comprising

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the steps of:

(1) growing a plant according to the invention which produces trehalose,

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- (2) harvesting the plant or plant parts which contain 5 trehalose, and
 - (3) air drying the plants or plant parts or alternatively,
 - (4) freeze drying the plants or plant parts.

The invention further comprises the plants and plant parts which have been preserved by a method according to the 10 invention.

The invention also includes a method for the production of trehalose comprising the steps of:

- (1) growing a plant which by virtue of a recombinant plant DNA genome is capable of producing (increased levels of) 15 trehalose,
 - (2) harvesting said plant or plant part,
 - (3) isolating the trehalose from the said plant or the said plant part.

The invention further includes a method for the production 20 of trehalose comprising the steps of:

- (1) growing in culture plant cells which by virtue of a recombinant plant DNA genome are capable of producing (increased levels of) trehalose,
- (2) isolating the trehalose from the said plant cell culture.
- 25 The invention further provides an isolated nucleic acid sequence encoding a trehalose phosphate synthase activity. A preferred isolated nucleic acid sequence is one obtained from E. coli, still more preferred is the isolated nucleic acid sequence represented in SEQIDNO: 2. Another preferred
- 30 embodiment comprises a nucleic acid sequence that codes for an amino acid sequence as in SEQIDNO: 3.

The following figures further illustrate the invention.

DESCRIPTION OF THE FIGURES.

35 Figure 1. Schematic representation of parts of the sucrose and starch biosynthetic pathways in plant sink tissues. The figure shows that carbohydrate produced in the leaf by photosynthesis is transported via the phloem tissue in the form of sucrose. Upon entering the sink it is unloaded by a

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membrane bound invertase activity to yield the monosugars glucose and fructose. By the action of a number of enzymatic steps these monosugars are converted to starch and/or sucrose as roughly shown here. The glucose metabolites G6P and UDPG are believed to be used as the substrates for the TPS-enzyme engineered into the plant by introduction of the plant expressible otsA gene. The figure shows how the amount of UDPG and G6P available as substrate is increased by reducing the levels of the enzymes SPS and AGPase. Their inhibition is marked with a cross.

	Figure 2.	Schematic map of the EBL4clone 7F11 from Kohara
		et al. (1987), containing the otsBA operon from
		E. coli. The 18.8 kb insert has been shaded.
15		•
13		The restriction sites for the enzymes EcoRV and
		<u>Hin</u> dIII used to clone the <u>ots</u> A gene are
		indicated, as well as their distance in kb with
		respect to the left-hand site of the insert.
		The <u>ots</u> A and B gene are indicated, the arrows
20		shows the direction of transcription. (See Fig
		11, extended map).
	Figure 3.	Schematic representation of binary vector
		pMOG663.
	Figure 4.	Sequence of the cloned potato SPS cDNA.
25		Underscore: maize SPS cDNA sequences used as
		oligonucleotides in the PCR amplification
		reaction.
	Figure 5.	Schematic representation of binary vector
		pMOG664.
30	Figure 6.	Schematic representation of binary vector
		pMOG665:
	Figure 7.	Schematic representation of binary vector
		pMOG666.
	Figure 8.	Restriction map of part of pTiB6 showing two
3 5		fragments cloned in pMOG579.
	Figure 9.	Schematic representation of pMOG579 used for
		constructing the helper plasmid without T-
		The state of the s

region in Agrobacterium strain MOG101.

Schematic representation of expression vector

Figure 10.

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pMOG180.

Figure 11. Nucleic acid sequence of the otsA gene and amino acid sequence of E. coli TPS.

Figure 12. Extended map of the EBL4clone 7F11 from Kohara et al. (1987), containing the otsBA operon from E. coli. The location of the TPS open reading frame (ORF) is indicated. (*: HindIII sites not present in the map of Kohara et al., infra)

Figure 13. Schematic representation of binary vector pMOG799.

Figure 14. Schematic representation of binary vector pMOG801.

Figure 15. Schematic representation of binary vector pMOG802.

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DETAILED DESCRIPTION OF THE INVENTION

A preferred embodiment of the invention comprises a potato plant capable of producing trehalose in tubers due to the presence in said potato plant of a plant expressible gene

20 which comprises in sequence:

- (a) a transcriptional initiation region derived from the 35S RNA of CaMV flanked upstream by a double enhancer,
- (b) a DNA sequence encoding trehalose phosphate synthase which is the coding region of the <u>ots</u>A gene located in the 25 <u>ots</u>BA operon of <u>E. coli</u>,
- (c) a transcriptional termination sequence derived from the nopaline synthase (nos) gene of Agrobacterium. Tubers of transgenic plants containing the plant expressible TPS gene produced trehalose, whereas control plants lacking this gene did not. Apparently, the trehalose phosphate which is produced by the transgenic tubers is converted into trehalose. Apparently, it is not required to provide for a trehalose phosphate phosphatase activity since it seems present in potato.
- Also illustrated in figure 1 is an approach to improve substrate availability for TPS. To this end two genes influencing the availability of glucose-6 phosphate (G6P) and UDPG, to whit an antisense SPS gene and a antisense APGase have been cloned under the control of the CaMV 35S promoter

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for expression in plant hosts. If introduced into a plant host containing a plant expressible TPS gene according to the invention, this will increase substrate availability for TPS and therefore trehalose synthesis. It will readily occur to someone skilled in the art that also other antisense genes may used to block the synthesis of sucrose or starch, in order to improve substrate availability.

Although the invention is described in detail for potato plants which express a plant expressible trehalose phosphate synthase gene from E. coli under the control of the CaMV 35S promoter as transcription initiation region, it will be clear to those of skill in the art that other spermatophytic plant hosts are equally suitable for the production of trehalose. Preferred plant hosts among the spermatophyta are the 15 Angiospermae, notably the Dicotyledoneae, comprising inter alia the Solanaceae as a representative family, and the Monocotyledoneae, comprising inter alia the Gramineae as a representative family. Suitable host plants, as defined in the context of the present invention include plants (as well 20 as parts and cells of said plants) and their progeny which have been genetically modified using recombinant DNA techniques to cause or enhance production of trehalose interest in the desired plant or plant organ; these plants may be used directly (e.g. the plant species which produce 25 edible parts) or after the trehalose is purified from said host (which be from edible as well as inedible plant hosts). Crops with edible parts according to the invention include those which have flowers such as cauliflower (Brassica oleracea), artichoke (Cynara scolymus), fruits such as apple 30 (Malus, e.g. domesticus), banana (Musa, e.g. acuminata), berries (such as the currant, Ribes, e.g. rubrum), cherries (such as the sweet cherry, Prunus, e.g. avium), cucumber (<u>Cucumis</u>, <u>e.g.</u> <u>sativus</u>), grape (<u>Vitis</u>, <u>e.g.</u> <u>vinifera</u>), lemon (Citrus limon), melon (Cucumis melo), nuts (such as the 35 walnut, <u>Juglans</u>, <u>e.g.</u> <u>regia</u>; peanut, <u>Arachis hypogeae</u>), orange (Citrus, e.g. maxima), peach (Prunus, e.g. persica), pear (Pyra, e.g. communis), pepper (Solanum, e.g. capsicum), plum (Prunus, e.q. domestica), strawberry (Fragaria, e.g.

moschata), tomato (Lycopersicon, e.g. esculentum), leafs,

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such as alfalfa (<u>Medicago</u>, <u>e.g. sativa</u>), cabbages (such as <u>Brassica oleracea</u>), endive (<u>Cichoreum</u>, <u>e.g. endivia</u>), leek (<u>Allium</u>, <u>e.g. porrum</u>), lettuce (<u>Lactuca</u>, <u>e.g. sativa</u>), spinach (<u>Spinacia e.g. oleraceae</u>), tobacco (<u>Nicotiana</u>, <u>e.g. g.g. sativa</u>)

5 tabacum), roots, such as arrowroot (Maranta, e.g. arundinacea), beet (Beta, e.g. vulgaris), carrot (Daucus, e.g. carota), cassava (Manihot, e.g. esculenta), turnip (Brassica, e.g. rapa), radish (Raphanus, e.g. sativus), yam (Dioscorea, e.g. esculenta), sweet potato (Ipomoea batatas)

10 and seeds, such as bean (<u>Phaseolus</u>, <u>e.g. vulgaris</u>), pea (<u>Pisum</u>, <u>e.g. sativum</u>), soybean (<u>Glycin</u>, <u>e.g. max</u>), wheat (<u>Triticum</u>, <u>e.g. aestivum</u>), barley (<u>Hordeum</u>, <u>e.g. vulgare</u>), corn (<u>Zea</u>, <u>e.g. mays</u>), rice (<u>Oryza</u>, <u>e.g. sativa</u>), tubers, such as kohlrabi (<u>Brassica</u>, <u>e.g. oleraceae</u>), potato (<u>Solanum</u>,

e.g. tuberosum), and the like. The edible parts may be conserved by drying in the presence of enhanced trehalose levels produced therein due to the presence of a plant expressible trehalose phosphate synthase construct. It may be advantageous to produce enhanced levels of trehalose, by

20 putting the DNA encoding the TPS activity under the control of an plant organ or tissue-specific promoter; the choice of which can readily be determined by those of skill in the art.

Any trehalose phosphate gene under the control of regulatory elements necessary for expression of DNA in plant cells, either specifically or constitutively, may be used, as long as it is capable of producing an active trehalose phosphate synthase activity. The nucleic acid sequence represented in SEQIDNO: 2, in fact any open reading frame encoding a trehalose phosphate synthase activity according to the invention, may be altered without necessarily altering the amino acid sequence of the protein encoded thereby. This fact is caused by the degeneracy of the genetic code. Thus the open reading frame encoding the trehalose phosphate synthase activity may be adapted to codon usage in the host plant of choice.

Also the isolated nucleic acid sequence represented by SEQIDNO: 2, may be used to identify trehalose phosphate synthase activities in other organisms and subsequently isolating them, by hybridising DNA from other sources with a

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DNA- or RNA fragment obtainable from the <u>E. coli</u> gene.

Preferably, such DNA sequences are screened by hybridising under stringent conditions (such as temperature and ionic strength of the hybridisation mixture. Whether or not conditions are stringent also depends on the nature of the hybridisation, <u>i.e.</u> DNA:DNA, DNA:RNA, RNA:RNA, as well as the length of the shortest hybridising fragment. Those of skill in the art are readily capable of establishing a stringent hybridisation regime.

Sources for isolating trehalose phosphate synthase activities include microorganisms (e.g. bacteria, yeast, fungi), plants, animals, and the like. Isolated DNA sequences encoding trehalose phosphate activity from other sources may be used likewise in a method for producing trehalose according to the invention.

The invention also encompasses nucleic acid sequences which have been obtained by modifying the nucleic acid sequence represented in SEQIDNO: 2 by mutating one or more codons so that it results in amino acid changes in the encoded protein, as long as mutation of the amino acid sequence does not entirely abolish trehalose phosphate synthase activity.

In principle any plant host is suitable in combination with any plant expressible trehalose phosphate synthase gene.

25 As trehalose genes from other sources become available these can be used in a similar way to obtain a plant expressible trehalose phosphate synthase gene combination as described here.

The inhibition of endogenous genes in order to enhance
30 substrate availability for the trehalose phosphate synthase,
as exemplified herein with the inhibition of endogenous
sucrose phosphate synthase gene and the ADP-Glucose
pyrophosphorylase gene, may be conducted in a number of ways
the choice of which is not critical to the invention.

Preferably gene inhibition is achieved through the so-called 'antisense approach'. Herein a DNA sequence is expressed which produces an RNA that is at least partially complementary to the RNA which encodes the enzymatic activity that is to be blocked (e.g. AGP-ase or SPS, in the examples).

It is preferred to use homologous antisense genes as these are more efficient than heterologous genes. The isolation of an antisense SPS gene from potato using a maize SPS-gene sequence as probe serves to illustrate the feasibility of this strategy. It is not meant to indicate that, for practicing the invention the use of homologous antisense fragments is required. An alternative method to block the synthesis of undesired enzymatic activities is the introduction into the genome of the plant host of an additional copy of an endogenous gene present in the plant host. It is often observed that such an additional copy of a gene silences the endogenous gene: this effect is referred to in the literature as the co-suppressive effect, or co-suppression.

15 In principle both dicotyledonous and monocotyledonous plants that are amenable for transformation, can be modified by introducing a plant expressible gene according to the invention into a recipient cell and growing a new plant that harbors and expresses the plant expressible gene. Preferred 20 plants according to the invention are those that are capable of converting trehalose-phosphate into trehalose, and which do contain no or little trehalose degrading activity. It will be understood that plants that lack the ability to convert the trehalose phosphate into trehalose are also included in 25 the present invention. These plants may be further modified by introducing additional genes that encode phosphatases that are capable of the conversion of trehalose phosphate into trehalose. In principle also plants are envisaged that do contain trehalases, since these plants can be made suitable 30 for the production of trehalose by inhibiting the activity of such enzymes, for instance by inhibiting expression of the genes encoding such enzymes using the antisense approach.

The method of introducing the plant expressible trehalosephosphate gene into a recipient plant cell is not crucial, as
long as the gene is stably incorporated into the genome of
said plant cell. In addition to the use of strains of the
genus <u>Agrobacterium</u> various other techniques are available
for the introduction of DNA into plant cells, such as
transformation of protoplasts using the calcium/polyethylene

glycol method, electroporation and microinjection or (coated) particle bombardment (Potrykus, 1990, Bio/Technol. 8, 535-542).

In addition to these so-called direct DNA transformation

5 methods, transformation systems involving vectors are widely
available, such as viral vectors (e.g. from the Cauliflower
Mosaic Virus (CaMV) and bacterial vectors (e.g. from the
genus Agrobacterium) (Potrykus, 1990, Bio/Technol. 8, 535542). After selection and/or screening, the protoplasts,

10 cells or plant parts that have been transformed can be
regenerated into whole plants, using methods known in the art
(Horsch et al., 1985, Science 225, 1229-1231).

It has been shown that monocotyledonous plants are amenable to transformation and that fertile transgenic plants 15 can be regenerated from transformed cells. The development of reproducible tissue culture systems for these crops, together with the powerful methods for introduction of genetic material into plant cells has facilitated transformation. Presently, preferred methods for transformation of monocots 20 are microprojectile bombardment of explants or suspension cells, and direct DNA uptake or electroporation (Shimamoto, et al, 1989, Nature 338, 274-276). Transgenic maize plants have been obtained by introducing the Streptomyces hygroscopicus bar-gene, which encodes phosphinothricin 25 acetyltransferase (an enzyme which inactivates the herbicide phosphinothricin), into embryogenic cells of a maize suspension culture by microprojectile bombardment (Gordon-Kamm, 1990, Plant Cell, 2, 603-618). The introduction of genetic material into aleurone protoplasts of other monocot 30 crops such as wheat and barley has been reported (Lee, 1989, Plant Mol. Biol. 13, 21-30). Wheat plants have been regenerated from embryogenic suspension culture by selecting only the aged compact and nodular embryogenic callus tissues for the establishment of the embryogenic suspension cultures 35 (Vasil, 1990 Bio/Technol. 8, 429-434). The combination with transformation systems for these crops enables the application of the present invention to monocots. These methods may also be applied for the transformation and regeneration of dicots.

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Monocotyledonous plants, including commercially important crops such as corn are amenable to DNA transfer by Agrobacterium strains (European patent 159 418 B1; Gould J, Michael D, Hasegawa O, Ulian EC, Peterson G, Smith RH, (1991) 5 Plant. Physiol. 95, 426-434).

As regards the choice of the host plant it is preferred to select plant species with little or no trehalose degrading activity. However, plants that do exhibit trehalase activity are not excluded from being a suitable host plant for the production of trehalose, although it may be necessary to provide for inhibition of trehalase activity if this prevents the accumulation of trehalose altogether. Such inhibition can be achieved using the antisense approach well known in the art, and illustrated for other purposes in this specification.

It should also be understood that the invention is not limited to the use of the CaMV 35S promoter as transcription initiation region. Suitable DNA sequences for control of expression of the plant expressible genes, including marker genes, such as transcriptional initiation regions, enhancers, non-transcribed leaders and the like, may be derived from any gene that is expressed in a plant cell which, such as endogenous plant genes, genes naturally expressed in plant cells such as those located on wild-type T-DNA of

Agrobacterium, genes of plant viruses, as well as other eukaryotic genes that include a transcription initiation region that conforms to the consensus sequence for eukaryotic transcription initiation. Also intended are hybrid promoters combining functional portions of various promoters, or synthetic equivalents thereof. Apart from constitutive promoters, inducible promoters, or promoters otherwise regulated in their expression pattern, e.g. developmentally or cell-type specific, may be used to control expression of the plant expressible genes according to the invention as long as they are expressed in plant parts that contain

To select or screen for transformed cells, it is preferred to include a marker gene linked to the plant expressible gene according to the invention to be transferred to a plant cell.

substrate for TPS.

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The choice of a suitable marker gene in plant transformation is well within the scope of the average skilled worker; some examples of routinely used marker genes are the neomycin phosphotransferase genes conferring resistance to kanamycin 5 (EP-B 131 623), the Glutathion-S-transferase gene from rat liver conferring resistance to glutathione derived herbicides (EP-A 256 223), glutamine synthetase conferring upon overexpression resistance to glutamine synthetase inhibitors such as phosphinothricin (WO87/05327), the acetyl transferase 10 gene from Streptomyces viridochromogenes conferring resistance to the selective agent phosphinothricin (EP-A 275 957), the gene encoding a 5-enolshikimate-3-phosphate synthase (EPSPS) conferring tolerance to Nphosphonomethylglycine, the bar gene conferring resistance against Bialaphos (e.g. W091/02071) and the like. The actual choice of the marker is not crucial as long as it is functional (i.e. selective) in combination with the plant cells of choice.

The marker gene and the gene of interest do not have to be linked, since co-transformation of unlinked genes (U.S. Patent 4,399,216) is also an efficient proces in plant transformation.

Preferred plant material for transformation, especially for dicotyledonous crops are leaf-discs which can be readily transformed and have good regenerative capability (Horsch R.B. et al., (1985) Science 227, 1229-1231).

Whereas the production of trehalose can be achieved with the plant expressible trehalose phosphate synthase gene as the sole carbohydrate modifying gene, the invention is further illustrated with examples of additional plant expressible antisense genes that are capable of effecting an increase of the availability of the substrate for trehalose phosphate synthase. Specific examples of such genes are the plant expressible antisense genes for SPS from maize and potato and AGPase from potato. The down regulation of carbohydrate modifying enzymes using the antisense approach is not limited by the specific examples. For instance partially complementary plant expressible antisense genes can be used to inhibit expression of a target gene, as long as

the plant expressible antisense gene produces a transcript that is sufficiently complementary with the transcript of the target gene and sufficiently long to inhibit expression said target gene.

- It is immaterial to the invention how the presence of two or more genes in the same plant is effected. This can inter alia done be achieved by one of the following methods:
 - (a) transformation of the plant line with a multigene construct containing more than one gene to be introduced,
- 10 (b) co-transforming different constructs to the same plant line simultaneously,
 - (c) subsequent rounds of transformation of the same plant with the genes to be introduced,
- (d) crossing two plants each of which contains a different gene to be introduced into the same plant.

The field of application of the invention lies both in agriculture and horticulture, for instance due to improved properties of the modified plants as such, as well as in any form of industry where trehalose is or will be applied.

- Trehalose phosphate and trehalose can be used as such for instance in purified form or in admixtures, or in the form of a storage product in plant parts. Plant parts harboring (increased levels of) trehalose phosphate or trehalose may be used as such or processed without the need to add trehalose.
- Also trehalose can be purified from the plants or plant parts producing it subsequently used in an industrial process. In the food industries trehalose can be employed by adding trehalose to foods before drying. Drying of foods is an important method of preservation in the industry.
- Trehalose seems especially useful to conserve food products through conventional air-drying, and to allow for fast reconstitution upon addition of water of a high quality product (Roser et al, July 1991, Trends in Food Science and Technology, pp. 166-169). The benefits include retention of
- natural flavors/fragrances, taste of fresh product, and nutritional value (proteins and vitamins). It has been shown that trehalose has the ability to stabilize proteins and membranes, and to form a chemically inert, stable glass. The low water activity of such thoroughly dried food products

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prevents chemical reactions, that could cause spoilage.

Field crops like corn, cassava, potato, sugar beet and sugarcane have since long been used as a natural source for bulk carbohydrate production (starches and sucrose). The 5 production of trehalose in such crops, facilitated by genetic engineering of the trehalose-biosynthetic pathway into these plant species, would allow the exploitation of such engineered crops for trehalose production.

All references cited in this specification are indicative 10 of the level of skill in the arts to which the invention pertains. All publications, whether patents or otherwise, referred to previously or later in this specification are herein incorporated by reference as if each of them was individually incorporated by reference.

15 The Examples given below are just given for purposes of enablement and do not intend in any way to limit the scope of the invention.

EXPERIMENTAL

20 DNA manipulations

All DNA procedures (DNA isolation from E.coli, restriction, ligation, transformation, etc.) are performed according to standard protocols (Sambrook et al. (1989) Molecular Cloning: a laboratory manual, 2nd ed. Cold Spring Harbor Laboratory 25 Press, CSH, New York).

Strains

In all examples E.coli K-12 strain DH5α is used for cloning. The Agrobacterium tumefaciens strain used for plant 30 transformation experiments is MOG101 which is a non-oncogenic octopine type helper strain derived form LBA1010 (Koekman et al. (1982) Plasmid 7, 119) by substitution of the T-DNA by a spectinomycin resistance marker.

35 Construction of Agrobacterium strain MOG101

A binary vector system (Hoekema A., Hirsch, P.R., Hooykaas, P.J.J., and Schilperoort, R.A. (1983) Nature 303, 179) is used to transfer gene constructs into potato plants. The helper plasmid conferring the Agrobacterium tumefaciens

virulence functions is derived from the octopine Ti-plasmid pTiB6. MOG101 is an Agrobacterium tumefaciens strain carrying a non-oncogenic Ti-plasmid (Koekman et al. 1982, supra) from which the entire T-region is deleted and substituted by a bacterial Spectinomycin resistance marker from transposon Tn1831 (Hooykaas et al., 1980 Plasmid 4, 64-75).

The Ti-plasmid pTiB6 contains two adjacent T-regions, TL (T-left) and TR (T-right). To obtain a derivative lacking the TL- and TR-regions, we constructed intermediate vector 10 pMOG579. Plasmid pMOG579 is a pBR322 derivative which contains 2 Ti-plasmid fragments homologous to the fragments located left and right outside the T-regions of pTiB6 (shaded in Figures 8 and 9). The 2 fragments are separated in pMOG579 by a 2.5 kb BamHI - HindIII fragment from transposon Tn1831 (Hooykaas et al., 1980 Plasmid 4, 64-75) carrying the 15 spectinomycin resistance marker (Figure 9). The plasmid is introduced into Agrobacterium tumefaciens strain LBA1010 [C58-C9 (pTiB6) = a cured C58 strain in which pTiB6 is introduced (Koekman et al. (1982), supra), by triparental 20 mating from E.coli, using HB101 8pRK2013 as a helper. Transconjugants are selected for resistance to Rifampicin (20 mg/l) and spectinomycin (250 mg/l). A double recombination between pMOG579 and pTiB6 resulted in loss of carbenicillin resistance (the pBR322 marker) and deletion of the entire T-25 region. Of 5000 spectinomycin resistant transconjugants replica plated onto carbenicillin (100 mg/l) 2 are found sensitive. Southern analysis (not shown) showed that a double

An alternative helper strain for MOG101 is <u>e.g.</u> LBA4404; this strain can also suitably be used for introduction of a binary plasmid, such as pMOG799 and subsequent plant transformation. Other suitable helper strains are readily available.

crossing over event had deleted the entire T-region. The resulting strain is called MOG101. This strain and its construction is analogous to strain GV2260 (Deblaere et al.

1985, Nucl. Acid Res. 13, 4777-4788).

Construction of the expression vector pMOG180

The expression vector pMOG180 is a derivative of pMOG18

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(EP 0 479 359 A1, Example 2b) wherein the gene coding for GUS is removed and other genes can be inserted between the AlMV RNA4 leader and 3' nos terminator as a BamHI fragment.

For this purpose, the EcoRI/NcoI fragment from pMOG18,

5 containing the 35S promoter and AlMV RNA4 leader sequences is synthesized using PCR technology with the primer sets 5'

GTTTCTACAGGACGGAGGATCCTGGAAGTATTTGAAAGA 3' and 5'

CAGCTATGACCATGATTACG 3' thus mutating the NcoI site into a BamHI site. pMOG18 vector is then cut with EcoRI and BamHI

10 after which the newly synthesized EcoRI and BamHI fragment can be ligated between these restriction sites. To circumvent PCR-induced random mutations in the promoter sequences, the EcoRI/EcoRV fragment in the PCR synthesized EcoRI/BamHI fragment is replaced by wildtype sequences from pMOG18. The short EcoRV/BamHI is checked for mutations by sequencing. The resulting expression vector is plasmid pMOG180 (Figure 10).

Triparental matings

The binary vectors pMOG663-666 are mobilized in triparental
matings with the <u>E. coli</u> strain HB101 containing plasmid
pRK2013 (Ditta G., Stanfield, S., Corbin, D., and Helinski,
D.R. et al. (1980) Proc. Natl. Acad. Sci. USA 77, 7347) into
Agrobacterium tumefaciens strain MOG101 and used for
transformation.

25

Transformation of potato

Potato (Solanum tuberosum cv. Désiree) is transformed with the Agrobacterium tumefaciens strain MOG101 containing the binary vector of interest as described (Hoekema A., Huisman, M.J., Molendijk, L., Van den Elzen, P.J.M., and Cornelissen, B.J.C. (1989) Bio/technology 7, 273). The basic culture medium is MS30R30, consisting of MS-medium (Murashige, T., and Skoog, F. (1962) Physiol. Plan. 14, 473), supplemented with 30 g/L sucrose, R3 vitamins (Ooms et al. G., Burrell, M.M., Karp, A., Bevan, M., and Hille, J. (1987) Theor. Appl. Genet. 73, 744), 5 µM zeatin riboside (ZR), and 0.3 µM indole acetic acid (IAA). The media are solidified where necessary, with 0.7 g/L Daichin agar.

Tubers of Solanum tuberosum cv. Désiree are peeled and

surface sterilized for 20 minutes in 0.6% hypochlorite solution containing 0.1% Tween-20. The potatoes are washed thoroughly in large volumes of sterile water for at least 2 hours. Discs of approximately 2 mm thickness are sliced from 5 cylinders of tuber tissue prepared with a corkbore. Discs are incubated for 20 minutes in a suspension consisting of the MS30R3 medium without ZR and IAA, containing 106-107 bacteria/ml of Agrobacterium MOG101 containing the binary vector. The discs are subsequently blotted dry on sterile 10 filter paper and transferred to solid MS30R3 medium with ZR and IAA. Discs are transferred to fresh medium with 100 mg/L cefotaxim and 50 mg/L vancomycin after 2 days. A week later, the discs are transferred again to the same medium, but this time with 100 mg/L kanamycin to select for transgenic shoots. 15 After 4-8 weeks, shoots emerging from the discs are excised and placed onto rooting medium (MS30R3-medium without ZR and IAA, but with 100 mg/L cefotaxim and 100 mg/L kanamycin). The shoots are propagated axenically by meristem cuttings and transferred to soil after root development. Where

20 appropriate, 10 mg/L hygromycin is used for selection instead of 100 mg/L kanamycin.

Trehalose assay

Trehalose is determined essentially as described by Hottiger 25 et al. (Hottiger et al. (1987) J. Bact. 169, 5518). Potato tuber tissue is frozen in liquid nitrogen, powdered with pestle and mortar and subsequently extracted for 60 minutes at room temperature in app. 3 volumes of 500 mM. trichloroacetic acid. After centrifugation the pellet is 30 extracted once more in the same way. The combined supernatants from the two extractions are assayed for anthrone positive material (Spiro R.G. (1966) Meth. Enzymol. 8, 3). Trehalose is determined qualitatively by TLC. The extracts are deionized (Merck, Ion exchanger V) and loaded 35 onto Silica Gel 60 plates (Merck). After chromatography plates are developed with n-butanol-pyridine-water (15:3:2, v/v). Spots are visualized by spraying with 5 mg/ml vanillin in concentrated H,SO, and heating at 130°C. Commercially available trehalose (Sigma) is used as a standard.

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Enzyme assays

In all determinations non-transgenic tuber material of variety Desiree is used as control. Protein content in all samples is determined as described by Bradford (Bradford (1976) Anal. Biochem. 72, 248). For assays on tuber extracts, frozen potato tuber slices of app. 100 mg are homogenized in 100 μ l 20 mM HEPES pH 7.4, centrifuged (Eppendorf, 5 minutes at maximum speed). The supernatant is used for activity assays.

10

TPS activity - TPS activity is determined essentially as described by Hottiger et al. (Hottiger T., Schmutz, P., and Wiemken, A. (1987) J. Bact. 169, 5518). Tuber extract assay mixtures contained 50 mM tricine (K') pH 7.0, 10 mM glucose-15 6-phosphate, 5mM UDP-glucose, 12.5 mM MgCl, in a total volume of 0.4 ml. In controls glucose-6-phosphate is omitted. Assay mixtures are incubated at 37°C for 5-30 min. The reaction is stopped by addition of 0.2 ml ice-cold 1 N perchloric acid. After neutralization with 0.2 ml 1 N KOH, 20 the samples are stored on ice for 10 minutes and subsequently centrifuged at 2,000 x g. UDP is determined in the supernatants. The assay mixture contained 140 mM tricine (Kt) pH 7.6, 2 mM phosphoenolpyruvate, 0.31 mM NADH, 20 U lactate dehydrogenase from rabbit muscle (Sigma Type XXXIX) in a 25 total volume of 1.96 ml. The reaction is started by addition of 20 U pyruvate kinase from rabbit muscle (Sigma Type III). The decrease of the absorbance at 340 nm at 370C is used to calculate the UDP concentration. One unit of TPS activity is defined as nmol UDP formed per min at 37°C.

30

AGPase activity - AGPase activity is determined as described by Müller-Röber et al. (Müller-Röber B., Sonnewald, U., and Willmitzer, L. (1992) EMBO J. 11, 1229). Production of glucose-1-phosphate from ADP-glucose is determined in a NAD-linked glucose-6-phosphate dehydrogenase system. The reaction assay contained 80 mM HEPES pH 7.4, 10 mM MgCl₂, 1 mM ADP-glucose, 0.6 mM NAD, 10 µM glucose-1,6-diphosphate, 3 mM DTT, 0.02% bovine serum albumin, 1 U phosphoglucomutase from rabbit muscle (Sigma), 2.5 U NAD-linked glucose-6-phosphate

dehydrogenase from Leuconostoc mesenteroides and tuber extract. The reaction is initiated by addition of sodiumpyrophosphate to a final concentration of 2 mM. NAD reduction is measured spectrophotometrically at 340 nm and 30°C. A unit of AGPase activity is defined as nmol glucose-1-phosphate generated per min at 30°C.

SPS activity - SPS activity is determined essentially as described by Lunn & ApRees (Lunn and ApRees (1990) Phytochem. 29, 1057). Assay mixtures contained 50 mM tricine (K*) pH 7.0, 5 mM fructose-6-phosphate, 5mM UDP-glucose, 12.5 mM MgCl,, tuber extract, and water in a total volume of 0.4 ml. In controls fructose-6-phosphate is omitted. Assay mixtures are incubated at 25°C for 5-30 min. The reaction is stopped by addition of 0.2 ml ice-cold 1 N perchloric acid. After neutralization with 0.2 ml 1 N KOH, the samples are stored on ice for 10 minutes and subsequently centrifuged at $2,000 \times g$. UDP is determined in the supernatants. The assay mixture contained 140 mM tricine (Kt) pH 7.6, 2 mM 20 phosphoenolpyruvate, 0.31 mM NADH, 20 U lactate dehydrogenase from rabbit muscle (Sigma Type XXXIX) in a total volume of 1.96 ml. The reaction is started by addition of 20 U pyruvate kinase from rabbit muscle (Sigma Type III). The decrease of the absorbance at 340 nm at 37°C is used to calculate the UDP

25 concentration. One unit of SPS activity is defined as nmole UDP formed per min at 37°C.

EXAMPLE I

Cloning of the Escherichia coli otsA gene

In E.coli trehalose phosphate synthase (TPS) is encoded by the otsA gene located in the operon otsBA. The location and the direction of transcription of this operon on the E.coli chromosome are precisely known (Kaassen I., Falkenberg, P., Styrvold, O.B., and Strom, A.R. (1992) J. Bact. 174, 889). It is located in the 41-42' region of the E.coli chromosome, and is confined on a 2.9 kb HindIII fragment on EMBL4 genomic clone designated 7F11 of the map by Kohara et al. (Kohara Y., Akiyama, K. and Isono, K. (1987) Cell 50, 495). The position of the otsBA operon on this clone 7F11 is shown in Figure 2.

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DNA is prepared from a lysate of lclone 7F11, and digested with HindIII. We isolated the 2.9 kb HindIII fragment containing otsBA (the 'righthand' HindII-site at 14.3 kb in the insert is omitted on the map by Kohara, as already 5 noticed by Kaassen). The 2.9 kb HindIII-fragment is cloned in pUC18 linearized with HindIII. From the resulting plasmid an EcoRV/HindIII fragment of 2.1 kb containing the otsA gene is isolated, it is made blunt using Klenow polymerase and then cloned in vector pMOG180 linearized with BamHI and made blunt 10 using Klenow polymerase. The resulting expression plasmid contained the E. coli otsA gene in the correct orientation under control of the Cauliflower Mosaic Virus (CaMV) 35S promoter with double enhancer (Guilley H., Dudley, R.K., Jonard, G., Balazs, E., and Richards, K.E. (1982) Cell 30, 763), the Alfalfa Mosaic Virus (AlMV) RNA4 leader sequence (Brederode et al. F.T., Koper-Zwarthoff, E.C., and Bol, J.F. (1980) Nucl. Acids Res. 8, 2213) and the nopaline synthase transcription terminator sequence from Agrobacterium tumefaciens. The expression cassette is cloned as an 20 EcoRI/HindIII fragment into the binary vector pMOG23 (deposited on January 29, 1990 at the Centraal Bureau voor Schimmelcultures under accession number 102.90) The resulting binary vector pMOG663 (see Figure 3) is used to transform potato.

25

Example II

Trehalose production in potato tubers transformed with pMOG663.

Potato tuber discs are transformed with the binary vector

pMOG663. Transgenic shoots are selected on kanamycin. A

number of 20 independent transgenic shoots containing the
plant expressible E.coli TPS-construct are analyzed for
trehalose phosphate synthase (TPS) activity. Shoots found to
contain the enzyme are grown to mature plants. Mature tubers

of those transgenic potato plants, analyzed for trehalose,
are found to contain elevated levels of trehalose in
comparison with non-transgenic control plants. Transgenic
plant line 663.1 is propagated for further work.

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Example III

Construction of pMOG664

Two oligonucleotides corresponding to the cDNA sequence of the small subunit of ADP-glucose pyrophosphorylase (AGPase) 5 from potato tuber (EMBL data bank accession number X61186) are synthesized. The sequences are as follows:

- 5' TCCCCATGGAATCAAAGCATCC 3' (SEQIDNO: 4)
- 5' GATTGGATCCAGGGCACGGCTG 3' (SEQIDNO: 5)
- 10 The oligonucleotides are designed to contain suitable restriction sites (BamHI and NcoI, underlined) at their termini to allow assembly in an expression cassette in an antisense orientation. A fragment of about 1 kb is PCR amplified with these oligonucleotides using DNA isolated from 15 a cDNA library from potato cv. Désiree prepared from 2 month old leaf tissue (Clontech) as a template. After sequencing it can be shown, that the fragment is identical with the AGPase sequence deposited in the EMBL data bank. Following digestion with BamHI and NcoI, the fragment is cloned in pMOG18 20 linearized with BamHI and NcoI. From the resulting plasmid the 1.85 kb EcoRI/BamHI fragment is isolated (containing the CaMV 35S promoter, the AlMV RNA4 leader and the AGPase fragment in an antisense orientation) as well as the 0.25 kb BamHI/HindIII fragment containing the nos-terminator. These 25 two fragments are cloned in a three-way ligation with the binary vector pMOG22 linearized with EcoRI and HindIII. The binary vector pMOG22 contains a plant expressible HPTII gene for hygromycin selection in transgenic plants (pMOG22 has been deposited at the Centraal Bureau voor Schimmelcultures 30 on January 29, 1990 under accession number 101.90). The resulting binary vector pMOG664 (see Figure 4) is used for potato transformation.

Example IV

35

Construction of pMOG665

A set of oligonucleotides complementary to the sequence of the maize sucrose phosphate synthase (SPS) cDNA (Worrell A.C., Bruneau, J-M., Summerfelt, K., Boersig, M., and Voelker, T.A. (1991) Plant Cell 3, 1121) is synthesized. WO 95/06126

Their sequences are as follows:

- 5' CTAGGTCGTGATTCTGATACAGGTGGCCAGGTG 3' (SEQIDNO: 6)
- 5' CAGCATCGGCATAGTGCCCATGTATCACGTAAGGC 3' (SEQIDNO: 7)

These oligonucleotides are used to PCR amplify a DNA fragment of 370 bp using DNA isolated from a potato cv. Désiree cDNA library prepared from 2 month old leaf tissue (Clontech) as a template. After sequencing of this fragment it can be shown that it is highly complementary to the SPS sequence of maize (see Figure 5, and Worrell et al. (1991) Plant Cell 3, 1121).

- The PCR amplified fragment is made blunt-ended and cloned in pMOG18 linearized with NcoI and BamHI and made blunt-ended with Klenow polymerase. From a clone with the SPS fragment in the antisense orientation with respect to the CaMV 35S promoter, the EcoRI/HindIII fragment is cloned into the
- binary vector pMOG22 linearized with EcoRI, in a three-way ligation using a synthetic adapter with the following sequence:
 - 5' AGCTTCCCCCCG 3' (SEQIDNO: 16)

;;;;;;;;

20 3' AGGGGGGGCTTAA 5' (SEQIDNO: 17)

The resulting binary vector pMOG665 (see Figure 6) is used for potato transformation.

25

35

Example IV

Construction of pMOG666

The EcoRI fragment of plasmid pMOG665 containing the antisense SPS cassette, is cloned in the binary vector pMOG664 (containing the antisense AGPase cassette) linearized with EcoRI. The resulting binary vector carrying the two anti-sense constructs is called pMOG666 (see Figure 7).

Example V

Trehalose production in potato transformed with pMOG663 and pMOG664

Potato tuber discs of kanamycin resistant transgenic plant line 663.1, expressing TPS (example II) are transformed with the binary vector pMOG664, containing the antisense AGPase construct. Transgenic shoots are selected on 10 mg/L

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hygromycin. Transgenic shoots are recovered, and checked by PCR for the presence of both pMOG663 and pMOG664 sequences. Transgenic plants containing the plant expressible E. coli TPS construct and the antisense AGPase construct are analyzed for TPS and AGPase activity.

Analysis of transgenic tubers for AGPase activity shows reductions in activity levels in individual transgenic lines in comparison with non-transgenic controls. Northern blotting shows that also mRNA levels for AGPase are reduced in the transgenic plants compared to those in non-transgenic control plants. Trehalose levels in tubers of transgenic potato plants, found to exhibit TPS activity, and having reduced levels of AGPase, show an increase in comparison with the

levels that can be found in tubers of transgenic plant line

15 663.1.

663.1.

Example VI

Trehalose production in potato transformed with pMOG663 and pMOG665

Potato tuber discs of transgenic plant line 663.1 expressing TPS are transformed with the binary vector pMOG665, containing the antisense SPS construct. Transgenic shoots are selected on 10 mg/L hygromycin. Emerging shoots are checked by PCR for the presence of both pMOG663 and pMOG665 sequences. Transgenic shoots containing the plant expressible E. coli TPS construct and the antisense SPS construct are

Analysis of transgenic tubers for SPS activity shows reductions in the levels for both enzymes in individual transgenic lines in comparison with non-transgenic controls. Northern blotting shows that also mRNA levels for SPS are reduced in the transgenic plants compared to those in non-transgenic control plants. Trehalose levels in tubers of transgenic potato plants, found to exhibit TPS activity, and having reduced levels of SPS, show an increase in comparison with the levels found in tubers of transgenic plant line

analyzed for TPS and SPS activity.

Example VII

Trehalose production in potato transformed with pMOG663 and pMOG666

5 Potato tuber discs of transgenic plant line 663.1 expressing TPS are transformed with the binary vector pMOG666, containing the two antisense AGPase and SPS constructs. Transgenic shoots are selected on 10 mg/L hygromycin. Emerging shoots are checked by PCR for the presence of the plant expressible E. coli TPS construct, and the antisense AGPase and SPS construct. Positive shoots are analyzed for TPS, AGPase and SPS activity.

Analysis of transgenic tubers for AGPase and SPS activity

15 shows reductions in the levels for both enzymes in individual transgenic lines in comparison with non-transgenic controls. Northern blotting shows that also mRNA levels are reduced in the transgenic plants compared to those in non-transgenic control plants. Trehalose levels in tubers of transgenic potato plants, found to exhibit TPS activity, and having reduced levels of AGPase and SPS, show an increase in comparison with the levels found in tubers of transgenic plant line 663.1.

25 The following examples describe the identification of the nucleotide sequence encoding a full length <u>E.coli</u> trehalose phosphate synthase activity. The amino acid sequence of the complete <u>E. coli</u> TPS is also disclosed.

30 Example VIII

Cloning of a full length E. coli otsA gene

In <u>E.coli</u> trehalose phosphate synthase (TPS) is encoded by the <u>ots</u>A gene located in the operon <u>ots</u>BA. The location and the direction of transcription of this operon on the <u>E.coli</u> chromosome are known (Kaasen, I., Falkenberg, P., Styrvold, O.B., and Ström, A.R. (1992) J. Bact. <u>174</u>, 889). The <u>ots</u>A gene is located at 42', and according to Kaasen et al. confined on a 18.8 kb fragment present in the EMBL4 genomic clone designated 7F11 of the map by Kohara et al. (Kohara,

Y., Akiyama, K., and Isono, K. (1987) Cell 50, 495). DNA prepared from a lysate of lambda clone 7F11, and digested with HindIII. The isolated 2.9 kb HindIII fragment (the 'right-hand' HindIII site at 14.3 kb in the insert was 5 omitted on the map by Kohara et al., as already noticed by Kaasen et al.) is cloned in pUC18 linearized with HindIII. The 2.9 kb HindIII insert from the resulting plasmid, designated pMOG674, is sequenced. The sequence is found to contain part of the araH gene of the arabinose transport 10 operon (Scripture, J.B., Voelker, C., Miller, S., O' Donnell, R.T., Polgar, L., Rade, J., Horazdovsky, B.F., and Hogg, R.W. (1987) J. Mol. Biol. 197, 37), the otsB gene encoding TPP as localized by Kaasen et al. and part of the otsA gene encoding TPS. The otsA is found not to be confined to the 2.9 kb 15 HindIII fragment as described by Kaasen et al. To complete the sequence an overlapping BamHI/EcoRI fragment is isolated and partially sequenced. The complete TPS-encoding sequence of the otsA gene is shown in Figure 11 (SEQIDNO: 2). The position of the otsA gene on clone 7F11, with the restriction 20 enzyme sites used, is shown in Figure 12. An additional HindIII site not present on the map published by Kohara et al. is found on the 'left-hand' site of the 2.9 kb HindIII fragment.

The HindIII site in pMOG180 is replaced by a SstI site, by cloning the oligonucleotide duplex:

```
SstI
5' AGCTCACGAGCTCTCAGG 3' (SEQIDNO: 8)
3' GTGCTCGAGAGTCCTCGA 5' (SEQIDNO: 9)
```

into pMOG180 cut with HindIII. The resulting vector is designated pMOG746. The oligonucleotide duplex:

```
BamHI SphI HindIII

SmaI | BamHI
| BamHI
| | SmaI | SamHI
```

is cloned in vector pMOG746 linearized with BamHI. The vector with the oligonucleotide duplex in the desired orientation (checked by restriction enzyme digestion) is designated

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pMOG747. The 2.9 kb HindIII fragment of plasmid pMOG674 is cloned in pMOG747 linearized with HindIII, resulting in vector pMOG748. The app. 2.4 kb EcoRV/SstI and the app. 3.5 kb SstI/SmaI fragments of pMOG748 are isolated, ligated and transformed into E. coli, thus deleting the 3' end of the 2.9 kb HindIII fragment. The resulting plasmid is designated pMOG749. The 5' end of the otsA gene is synthesized by PCR using the synthetic oligonucleotides TPS1 and TPS2 with pMOG749 as a template.

10

TPS1 5' GAGAAAATACCCGGGGTGATGAC 3' (SEQIDNO: 12)

TPS2 5' GATAATCGTGGATCCAGATAATGTC 3' (SEQIDNO: 13)

By sequencing it is confirmed that the 0.4 kb PCR fragment

15 has the correct sequence. The 1 kb BamHI/HindIII fragment of

pMOG749 is cloned together with the 0.4 kb XmaI/BamHI PCR

fragment in pMOG747 linearized with XmaI and HindIII. In the

resulting plasmid, digested with HindIII and SstI, the

synthetic oligonucleotide duplex TPS6/7 is cloned, encoding

20 the three C-terminal amino acids of TPS.

LysLeuAlaStop

5' AGCTGGCGTGAGGAGCGGTTAATAAGCTTGAGCT 3' (SEQIDNO: 14)

3' CCGCACTCCTCGCCAATTATTCGAAC 5' (SEQIDNO: 15)

25

In the resulting plasmid, digested with HindIII and SstI, the 0.25 kb HindIII/SstI fragment of plasmid pMOG749 is cloned, comprising the terminator from the Agrobacterium tumefaciens nopaline synthase (NOS) gene, resulting in plasmid pMOG798. 30 This plasmid contains the E. coli otsA gene in the correct orientation under control of the Cauliflower Mosaic Virus (CaMV) 35S promoter with double enhancer (Guilley et al. (1982) Cell 30, 763), the Alfalfa Mosaic Virus (AMV) RNA4 leader sequence (Brederode et al. (1980) Nucl. Acids Res. 8, 35 2213) and the nopaline synthase transcription terminator sequence from Agrobacterium tumefaciens. The expression cassette is cloned as a 2.5 kb EcoRI/SstI fragment into the binary vector pMOG23 linearized with EcoRI and SstI. The resulting binary vector, pMOG799 (Fig. 13), is used to

- 30 -

transform potato (An <u>E. coli</u> strain harbouring pMOG799 has been deposited at the Centraal Bureau voor Schimmelcultures, Phabagen collections, Padualaan 8, Utrecht, The Netherlands, on August 23, 1993, deposit number CBS 430.93).

5 .

Example IX

Potato tuber discs are transformed with the binary vector pMOG799 using Agrobacterium tumefaciens. Transgenic shoots are selected on kanamycin. A number of 20 independent transgenic shoots are analyzed for trehalose phosphate synthase (TPS) activity. Shoots found to contain the enzyme are grown to mature plants. Analyses of mature tubers of those transgenic potato plants show elevated levels of trehalose in comparison with non-transgenic control plants. Transgenic plant line MOG799.1 is propagated for further work.

Example X

Construction of pMOG664

Two oligonucleotides corresponding to the cDNA sequence of the small subunit of ADP-glucose pyrophosphorylase (AGPaseB) from potato tuber cv. Désirée (Müller-Röber, B., Kossmann, J., Hannah, L.C., Willmitzer, L., and Sonnewald, U. (1990) Mol. Gen. Genet. 224, 136-146) are synthesized:

25,

- 5' TCCCCATGGAATCAAAGCATCC 3' (SEQIDNO: 4)
- 5' GATTGGATCCAGGGCACGGCTG 3' (SEQIDNO: 5)

The oligonucleotides are designed to contain suitable restriction sites (BamHI and NcoI, underlined) at their termini to allow assembly in an expression cassette in an antisense orientation after digestion with these enzymes. A fragment of about 1 kb is PCR amplified with these oligonucleotides using DNA isolated from a cDNA library from potato cv. Désiree prepared from 2 month old leaf tissue (Clontech) as a template. By sequencing it is shown, that the fragment is identical with the AGPase B sequence from potato cv. Désirée (Müller-Röber, B., Kossmann, J., Hannah, L.C., Willmitzer, L., and Sonnewald, U. (1990) Mol. Gen. Genet.

- 31 -

224, 136-146). Following digestion with BamHI and NcoI, the fragment is cloned in pMOG18 linearized with BamHI and NcoI. From the resulting plasmid the 1.85 kb EcoRI/BamHI fragment (containing the CaMV 35S promoter, the AMV RNA4 leader and the AGPase fragment in an antisense orientation), as well as the BamHI/HindIII fragment containing the terminator from the nopaline synthase (NOS) gene from Agrobacterium tumefaciens are cloned in a three-way ligation in the binary vector pMOG22 linearized with EcoRI and HindIII. The binary vector pMOG22 contains a plant expressible HPTII gene for hygromycin selection in transgenic plants (pMOG22 has been deposited at the Centraal Bureau voor Schimmelcultures on January 29, 1990 under accession number 101.90). The resulting binary vector pMOG664 (Fig. 4) is used for potato transformation.

15

Example XI

Construction of pMOG801

A set of oligonucleotides complementary to the sequence of the maize sucrose phosphate synthase (SPS) cDNA (Worrell, O A.C., Bruneau, J-M., Summerfalt, K., Boersig, M., and Voelker, T.A. (1991) Plant Cell 3, 1121) is synthesized. Their sequences are as follows:

- 5' CTAGGTCGTGATTCTGATACAGGTGGCCAGGTG 3' (SEQIDNO: 6)
- 25 5' CAGCATCGGCATAGTGCCCATGTATCACGTAAGGC 3' (SEQIDNO: 7)

These oligonucleotides are used to PCR amplify a DNA fragment of 370 bp using DNA isolated from a potato cv. Désiree cDNA library prepared from 2 month old leaf tissue (Clontech) as a template. By sequencing of this fragment it is shown, that it is homologous to the SPS sequence of maize (see Figure 4, and Worrell et al. (1991). The PCR fragment is used to screen a lambda gt10 library of potato cv. Désiree cDNA library prepared from 2 month old leaf tissue (Clontech). The insert of one positively hybridizing clone is sequenced. The sequence of the 654 bp DNA fragment is found to be 65% identical with the corresponding part of the maize SPS sequence (Starting at nucleotide number 349 in Figure 11 in Worrell et al. (1991). The EcoRI insert of this clone is

- 32 -

cloned in pMOG180 digested with BamHI, in a three-way ligation with the following synthetic oligonuclotide duplex.

- 5' GATCGTCAGATCTAGC 3' (SEQIDNO: 14)
- 5 3' CAGTCTAGATCGTTAA 5' (SEQIDNO: 15)

The plasmid, having the SPS fragment in the antisense orientation with respect to the CaMV 35S promoter, is designated pMOG787. The EcoRI/HindIII fragment of plasmid 10 pMOG787 is cloned in a three-way ligation with a synthetic linker:

- 5' AGCTTCCCCCCG 3' (SEQIDNO: 16)
- 3' AGGGGGGCTTAA 5' (SEQIDNO: 17)

15

into the binary vector pMOG22 linearized with EcoRI. The binary vector pMOG22 contains a plant expressible HPTII gene for hygromycin selection in transgenic plants (pMOG22 has been deposited at the Centraal Bureau voor Schimmelcultures on January 29, 1990 under accession number 101.90). The resulting binary vector pMOG801 (Fig. 14) is used for potato transformation.

Example XII

25

Construction of pMOG802

The EcoRI fragment of plasmid pMOG801, containing the antisense SPS expression cassette, is cloned in the binary vector pMOG664 (containing the antisense AGPase cassette), linearized with EcoRI. The resulting binary vector is called pMOG802 (Fig 15).

Example XIII

Trehalose production in potato transformed with pMOG799 and pMOG664

35 Potato tuber discs of kanamycin resistant plant line MOG799.1, expressing TPS (Example IX) are transformed with the binary vector pMOG664, containing the antisense AGPase expression cassette. Transgenic shoots, selected on 10 mg/L hygromycin, are analyzed for the presence of the TPS and

- 33 -

antisense AGPase sequences by PCR. Transgenic plants containing both are analyzed for TPS and AGPase activity.

By analysis of transgenic tubers for AGPase activity it is shown that, reductions in activity levels in individual transgenic lines in comparison with non-transgenic controls occur. By Northern blots it is shown, that mRNA levels for AGPase are reduced in the transgenic plants compared to those in non-transgenic control plants. Trehalose levels in tubers of transgenic potato plants, found to exhibit TPS activity, and having reduced levels of AGPase, show an increase in comparison with the levels found in tubers of transgenic plant line MOG799.1.

Example XIV

Trehalose production in potato transformed with pMOG799 and pMOG801

Potato tuber discs of kanamycin resistant plant line MOG799.1, expressing TPS (Example IX) are transformed with the binary vector pMOG801, containing the antisense SPS expression cassette. Transgenic shoots, selected on 10 mg/L hygromycin, are analyzed for the presence of the TPS and antisense SPS sequences by PCR. Transgenic plants containing both are analyzed for TPS and SPS activity.

By analysis of transgenic tubers for SPS activity it is shown
that reductions in activity levels in individual transgenic
lines in comparison with non-transgenic controls occur. By
Northern blots it is shown, that mRNA levels for SPS are
reduced in the transgenic plants compared to those in nontransgenic control plants. Trehalose levels in tubers of
transgenic potato plants, found to exhibit TPS activity, and
having reduced levels of SPS, show an increase in comparison
with the levels found in tubers of transgenic plant line
MOG799.1.

35 Example XV

Trehalose production in potato transformed with pMOG799 and pMOG802

Potato tuber discs of kanamycin resistant plant line MOG799.1, expressing TPS (Example IX) are transformed with

- 34 -

the binary vector pMOG802, containing the antisense SPS and AGPase expression cassettes. Transgenic shoots, selected on 10 mg/L hygromycin, are analyzed for the presence of the TPS, antisense AGPase and antisense SPS sequences by PCR.

5 Transgenic plants containing all three constructs are analyzed for TPS, AGPase and SPS activity.

By analysis of transgenic tubers for AGPase and SPS activity it is shown, that reductions in the activity levels for both enzymes in individual transgenic lines in comparison with 10 non-transgenic controls occur. By Northern blots it is shown that mRNA levels for AGPase and SPS are reduced in the transgenic plants compared to those in non-transgenic control plants. Trehalose levels in tubers of transgenic potato plants, found to exhibit TPS activity, and having reduced levels of SPS, show an increase in comparison with the levels found in tubers of transgenic plant line MOG799.1.

- 35 -

SEQUENCE LISTING

5	(1) GENERAL INFORMATION:	
5	(i) APPLICANT:	·
	(A) NAME: MOGEN International N.V. (B) STREET: Einsteinweg 97	
	(C) CITY: LEIDEN	
LO	(D) STATE: Zuid-Holland	
	(E) COUNTRY: The Netherlands (F) POSTAL CODE (ZIP): NL-2333 CB	
	(G) TELEPHONE: (31).(71).258282	
_	(H) TELEFAX: (31).(71).221471	
15	(ii) TITLE OF INVENTION: PRODUCTION OF TREHALOSE IN PLANTS	
	(iii) NUMBER OF SEQUENCES: 17	
20	(iv) COMPUTER READABLE FORM:	
	(A) MEDIUM TYPE: Floppy disk	
	(B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS	
	(D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)	
25		
	(V) CURRENT APPLICATION DATA: APPLICATION NUMBER: WO PCI/EP93/02290	
	101010101010101010100000000000000000000	
30	(2) INFORMATION FOR SEQ ID NO: 1:	
30	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGIH: 370 base pairs	
	(B) TYPE: nucleic acid (C) STRANDEDNESS: double	
35	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA to mRNA	
	(iii) HYPOIHETICAL: NO	
10	(III) mromentan. No	
	(vi) ORIGINAL SOURCE:	
	(A) ORGANISM: Solanum tuberosum (B) STRAIN: Desiree	
	(F) TISSUE TYPE: Leaf	
45	• •	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:	·
50	CIAGGIOGIG ATTCIGATAC AGGIOGOCAG GIGAAGIAIG TAGTAGAGCT TGCTOGAGCA	60
	CITGCAAACA TGAAAGGAGT TCACOGAGIT GATCTCTTGA CTOGGCAGAT CACATCCCCA	120
	GAGGITGATT CTAGCTATCG TGAGCCAATT GAGATGCTCT CATGCCCCATC TGATGCTTTG	180
55	CONCONCINCE DECOMPACIONE DOSCARGOSTI COCCAGONACE DISACARACADA DIDECARACADA ANCA	240

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	ATTTACATAC CAGAATTTGT TGATGGAGCA TTAAGCCACA TTGTGAATAT GGCAAGGGCT	300
	ATAGGGGAGC AAGTCAATGC TGGAAAAGCA GTGTGGCCTT ACGTGATACA TGGGCACTAT	360
5	GCCGATGCTG	370
	(2) INFORMATION FOR SEQ ID NO: 2:	
10	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1446 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear 	
15	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOIHETICAL: NO	
20	(vi) ORIGINAL SOURCE: (A) ORGANISM: Escherichia coli	
	(vii) IMMEDIATE SOURCE: (B) CLONE: 7F11	
25	(viii) Position in Genome: (B) Map Position: 41-42'	
30	<pre>(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 191446 (D) OTHER INFORMATION: /product= "trehalose phosphate synthase"</pre>	
35	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:	
	GAGAAAATAA CAGGAGTG ATG ACT ATG AGT CGT TTA GTC GTA GTA TCT AAC	51
40	Met Thr Met Ser Arg Leu Val Val Ser Asn 1 5 10	01
	CGG ATT GCA CCA CCA GAC GAG CAC GCC GCC AGT GCC GGT GGC CTT GCC Arg Ile Ala Pro Pro Asp Glu His Ala Ala Ser Ala Gly Gly Leu Ala 15 20 25	99
45	GTT GGC ATA CTG GGG GCA CTG AAA GCC GCA GGC GGA CTG TGG TTT GGC Val Gly Ile Leu Gly Ala Leu Lys Ala Ala Gly Gly Leu Trp Phe Gly 30 35 40	147
50	TGG AGT GGT GAA ACA GGG AAT GAG GAT CAG COG CTA AAA AAG GTG AAA Trp Ser Gly Glu Thr Gly Asn Glu Asp Gln Pro Leu Lys Lys Val Lys 45 50 55	195
55	AAA GGT AAC ATT ACG TGG GCC TCT TTT AAC CTC AGC GAA CAG GAC CTT Lys Gly Asn Ile Thr Trp Ala Ser Phe Asn Leu Ser Glu Gln Asp Leu 60 65 70 75	243

- 37 -

							CTC Leu			291
5							CCT Pro			339
10							TTA Leu			387
1 5	CAA Gln						CAC His 135			435
20							OGC Arg			483
		_					AAC Asn			531
25							GAT Asp			579
30							TGT Cys			627
35							ACA Thr 215			675
40							GAA Glu			723
			Ala				CIG Leu			771
45							GIC Val			819
50							TAT Tyr			867
55							TAT Tyr 295			915

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						CAG Gln 310			Ġ	963
5		 	 	 	 	 aaa Lys	 	 	 10	011
10						TTT Phe			10	059
1 5		 		 		TTA Leu		 	 1.	107
20						GIT Val			1	155
						TTT Phe 390			12	203
25						TAC Tyr			12	251
30						TCG Ser			13	299
35	TCC Ser					GTG Val			1:	347
40						aag Lys			13	395
40						GCT Ala 470			14	443
4 5	GOG Ala								14	446

50 (2) INFORMATION FOR SEQ ID NO: 3:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGIH: 476 amino acids
(B) TYPE: amino acid
(D) TOPOLOGY: linear

- 39 -

(ii) MOLECULE TYPE: protei	(ii)	MOLECULE	TYPE:	protein
----------------------------	------	----------	-------	---------

16:01	CENTENTOS	DESCRIPTION:	CEN	TD NO.	2.
(XI)	SECULNCE	DESCRIPTION:	SEU	TO NO:	3:

5 Met Thr Met Ser Arg Leu Val Val Val Ser Asn Arg Ile Ala Pro Pro 1 5 10 15

Asp Glu His Ala Ala Ser Ala Gly Gly Leu Ala Val Gly Ile Leu Gly
20 25 30

10
Ala Leu Lys Ala Ala Gly Gly Leu Trp Phe Gly Trp Ser Gly Glu Thr
35
40
45

Gly Asn Glu Asp Gln Pro Leu Lys Lys Val Lys Lys Gly Asn Ile Thr 15 50 55 60

Trp Ala Ser Phe Asn Leu Ser Glu Gln Asp Leu Asp Glu Tyr Tyr Asn 65 70 75 80

20 Gln Phe Ser Asn Ala Val Leu Trp Pro Ala Phe His Tyr Arg Leu Asp 85 90 95

Leu Val Gln Phe Gln Arg Pro Ala Trp Asp Gly Tyr Leu Arg Val Asn 100 105 110

25
Ala Leu Leu Ala Asp Lys Leu Leu Pro Leu Leu Gln Asp Asp Asp Ile
115
120
125

Ile Trp Ile His Asp Tyr His Leu Leu Pro Phe Ala His Glu Leu Arg 30 130 135 140

Lys Arg Gly Val Asn Asn Arg Ile Gly Phe Phe Leu His Ile Pro Phe 145 150 155 160

35 Pro Thr Pro Glu Ile Phe Asn Ala Leu Pro Thr Tyr Asp Thr Leu Leu 165 170 175

Glu Gln Leu Cys Asp Tyr Asp Leu Leu Gly Phe Gln Thr Glu Asn Asp 180 185 190

Arg Leu Ala Phe Leu Asp Cys Leu Ser Asn Leu Thr Arg Val Thr Thr 195 200 205

40

55

Arg Ser Ala Lys Ser His Thr Ala Trp Gly Lys Ala Phe Arg Thr Glu 45 210 215 220

Val Tyr Pro Ile Gly Ile Glu Pro Lys Glu Ile Ala Lys Gln Ala Ala 225 230 235 240

50 Gly Pro Leu Pro Pro Lys Leu Ala Gln Leu Lys Ala Glu Leu Lys Asn 245 250 255

Val Gln Asn Ile Phe Ser Val Glu Arg Leu Asp Tyr Ser Lys Gly Leu 260 265 270

Pro Glu Arg Phe Leu Ala Tyr Glu Ala Leu Leu Glu Lys Tyr Pro Gln

SUBSTITUTE SHEET

- 40 -

275 280 285 His His Gly Lys Ile Arg Tyr Thr Gln Ile Ala Pro Thr Ser Arg Gly 295 5 Asp Val Gln Ala Tyr Gln Asp Ile Arg His Gln Leu Glu Asn Glu Ala Gly Arg Ile Asn Gly Lys Tyr Gly Gln Leu Gly Trp Thr Pro Leu Tyr 10 330 Tyr Leu Asn Gln His Phe Asp Arg Lys Leu Leu Met Lys Ile Phe Arg Tyr Ser Asp Val Gly Leu Val Thr Pro Leu Arg Asp Gly Met Asn Leu Val Ala Ilys Glu Tyr Val Ala Ala Gln Asp Pro Ala Asn Pro Gly Val 20 Leu Val Leu Ser Gln Phe Ala Gly Ala Ala Asn Glu Leu Thr Ser Ala 390 Leu Ile Val Asn Pro Tyr Asp Arg Asp Glu Val Ala Ala Ala Leu Asp 25 410 Arg Ala Leu Thr Met Ser Leu Ala Glu Arg Ile Ser Arg His Ala Glu Met Leu Asp Val Ile Val Lys Asn Asp Ile Asn His Trp Gln Glu Cys Phe Ile Ser Asp Leu Lys Gln Ile Val Pro Arg Ser Ala Glu Ser Gln 450 455 35 Gln Arg Asp Lys Val Ala Thr Phe Pro Lys Leu Ala 470 (2) INFORMATION FOR SEQ ID NO: 4: 40 (i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single 45 (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA (iii) HYPOTHETICAL: YES 50

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

55 TCCCCATGGA ATCAAAGCAT CC

W	n	95	ME	126

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	(2) INFORMATION FOR SEQ ID NO: 5:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(iii) MOLECULE TYPE: cDNA (iii) HYPOTHETICAL: YES	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5: GATTGGATCC AGGGCACGGC TG	22
		22
20	(2) INFORMATION FOR SEQ ID NO: 6:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) IENGIH: 33 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: CDNA	
30	(iii) HYPOTHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
35	CTAGGTOGTG ATTCTGATAC AGGTGGCCAG GTG	33
	(2) INFORMATION FOR SEQ ID NO: 7:	
40	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
45	(ii) MOLECULE TYPE: DNA (genomic)	
	(iii) HYPOIHETICAL: YES	•
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
	CAGCATOGGC ATAGTGCCCCA TGTATCACGT AAGGC	35
55	(2) INFORMATION FOR SEQ ID NO: 8:	
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5	(i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
10	(iii) HYPOIHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:	
15	ACCICACGAG CICICAGG	18
	(2) INFORMATION FOR SEQ ID NO: 9:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 18 base pairs (B) TYPE: mucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: cDNA	
	(iii) HYPOIHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9: GIGCTOGAGA GICCTOGA	1 8
35	(2) INFORMATION FOR SEQ ID NO: 10:	•
40	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	
45	(iii) HYPOIHETICAL: YES	
50	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10: GATCCCCCCG GGCATGCAAG CITG (2) INFORMATION FOR SEQ ID NO: 11:	24
55	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGIH: 24 base pairs	

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	(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
5	(ii) MOLECULE TYPE: cDNA	
	(iii) HYPOIHETICAL: YES	
10	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
	GGGGCCCCGT ACGITCGAAC CTAG	24
15	(2) INFORMATION FOR SEQ ID NO: 12:	
20	(i) SEQUENCE CHARACTERISTICS: (A) IENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA	
25	(iii) HYPOTHETICAL: YES	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	
	GAGAAAATAC COGGGGTGAT GAC	23
	(2) INFORMATION FOR SEQ ID NO: 13:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 25 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
40	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
	(iii) HYPOIHETICAL: YES	
45	·	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:	
50	GATAATOGIG GATCCAGATA ATGIC	25
	(2) INFORMATION FOR SEQ ID NO: 14:	
55	(i) SEQUENCE CHARACTERISTICS: (A) LENGIH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEINESS: single	

- 44

	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: CDNA	
5	(iii) HYPOIHETICAL: YES	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
10	GATOSTCAGA TCTAGC	16
	(2) INFORMATION FOR SEQ ID NO: 15:	
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20	(ii) MOLECULE TYPE: CDNA	
	(iii) HYPOIHETICAL: YES	•
25	(<i>)</i>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	
	CAGICIAGAT OGITAA	16
30	(2) INFORMATION FOR SEQ ID NO: 16:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 13 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: CDNA	
40	(iii) HYPOTHETTCAL: YES	•
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:	
	ACCITCCCCC CCG	13
50	(2) INFORMATION FOR SEQ ID NO: 17:	
55	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 13 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

- 45 -

(ii) MOLECULE TYPE: cDNA

(iii) HYPOIHETTCAL: YES

5

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

AGGGGGGCT TAA

13

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CLAIMS

1. A plant expressible gene which when expressed in a plant or plant cell increases the trehalose content of said plant or plant cell.

- 2. A plant expressible gene according to claim 1 which comprises in sequence:
- (a) a transcriptional initiation region that is functional in said plant host,
- 10 (b) a DNA sequence encoding a trehalose phosphate synthase activity, and optionally
 - (c) a transcriptional termination sequence that is functional in said plant host.
- 15 3. A DNA sequence containing a plant expressible gene which comprises in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
- (b) a DNA sequence encoding a trehalose phosphate synthase 20 activity, and optionally
 - (c) a transcriptional termination sequence that is functional in said plant host,
 - and a plant expressible gene comprising in sequence:
- (a) a transcriptional initiation region that is functional in25 said plant host,
 - (b) a DNA sequence encoding an RNA sequence at least partially complementary to an RNA sequence which encodes sucrose phosphate synthase enzyme (SPS) naturally occurring in said plant host, and optionally
- 30 (c) a transcriptional termination sequence that is functional in said plant host.
 - 4. A DNA sequence comprising a plant expressible gene which comprises in sequence:
- 35 (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding a trehalose phosphate synthase

activity, and optionally

(c) a transcriptional termination sequence that is functional in said plant host, and

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- a plant expressible gene comprising in sequence:
- 5 (a) a transcriptional initiation region that is functional in said plant host.
- (b) a DNA sequence encoding an RNA sequence at least partially complementary to an RNA sequence which encodes a ADP-glucose pyrophosphorylase enzyme naturally occurring in 10 said plant host, and optionally
 - (c) a transcriptional termination sequence that is functional in said plant host.
- A DNA sequence comprising a plant expressible 15 gene which comprises in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding a trehalose phosphate synthase activity, and optionally
- 20 (c) a transcriptional termination sequence that is functional in said plant host,
 - and a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
- 25 (b) a DNA sequence encoding an RNA sequence at least partially complementary to an RNA sequence which encodes a sucrose phosphate synthase enzyme naturally occurring in said plant host, and optionally
- (c) a transcriptional termination sequence that is functional 30 in said plant host,
 - and a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
- a DNA sequence encoding an RNA sequence at least 35 partially complementary to an RNA sequence which encodes a ADP-glucose pyrophosphorylase enzyme naturally occurring in sai plant host, and optionally

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- (c) a transcriptional termination sequence that is functional in said plant host.
- 6. A vector suitable for cloning which comprises a 5 plant expressible gene according to claim 1 or 2.
 - 7. A vector suitable for cloning which comprises a DNA sequence of any one of the claims 3 to 5.
- 10 8. A vector according to claim 6 or 7 which is a binary vector.
 - 9. A microorganism comprising a vector of any one of the claims 6 to 8.

15

- 10. The microorganism of claim 9 which is of the genus Agrobacterium.
- A method for obtaining a plant capable of producing
 trehalose comprising the steps of,
 - (1) introducing into a recipient cell of a plant a plant expressible gene which when expressed in a plant or plant cell increases the trehalose content of said plant or plant cell,
- 25 and a plant expressible gene comprising in sequence:
 - (a) a transcriptional initiation region that is functional in said plant host,
 - (b) a DNA sequence encoding a selectable marker gene that is functional in said plant host, and optionally
- 30 (c) a transcriptional termination sequence that is functional in said plant host,
 - (2) generating a plant from a transformed cell under conditions that allow for selection for the presence of the selectable marker gene.

35

12. A recombinant plant DNA genome which contains a plant expressible trehalose phosphate synthase gene that is

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not naturally present therein.

- 13. A recombinant plant DNA genome which comprises
- (a) a plant expressible gene encoding trehalose phosphate5 synthase, and
 - (b) a plant expressible gene capable of inhibiting the biosynthesis of a sucrose phosphate synthesis activity.
 - 14. A recombinant plant DNA genome which comprises:
- 10 (a) a plant expressible gene encoding trehalose phosphate synthase,
 - (b) a plant expressible gene capable of inhibiting the biosynthesis of an ADP-Glucose pyrophosphorylase activity.
- 15 15. A recombinant plant DNA genome which comprises:
 - (a) a plant expressible gene encoding trehalose phosphate synthase,
- (b) a plant expressible gene capable of inhibiting the biosynthesis of an ADP-Glucose pyrophosphorylase activity, 20 and
 - (c) a plant expressible gene capable of inhibiting the biosynthesis of an sucrose phosphate synthesis activity.
- 16. A plant cell having a recombinant plant DNA genome 25 of any one of the claims 12 to 15.
 - 17. The plant cell of claim 16 which contains increased levels of trehalose compared with a plant cell of the same species having a non-recombinant plant DNA genome.
 - 18. A plant cell culture comprising plant cells of any one of the claims 16 or 17.
- 19. A method for the production of trehalose comprising 35 the steps of:
 - (1) growing in culture plant cells which by virtue of a recombinant plant DNA genome are capable of producing

- 50 -

- (increased levels of) trehalose,
- (2) isolating the trehalose from the said plant cell culture.
- 20. The method of claim 19 wherein the plant cell 5 culture is that of claim 18.
 - 21. A plant containing a cell of any one of the claims 16 to 17.
- 10 22. A plant consisting predominantly of cells of any one of the claims 16 to 17.
 - 23. A plant capable of producing increased levels of trehalose as a result of genetic modification.

15

- 24. A plant having a recombinant plant DNA genome of any one of the claims 13 to 15.
- 25. The plant of any one of the claims 23 to 24 which 20 contains increased levels of trehalose.
 - 26. The plant of claim 25 which is belongs to the Angiospermae.
- 25 27. A part of a plant containing a cell of any one of the claims 16 to 17.
 - 28. A part of a plant consisting predominantly of a cell of any one of the claims 16 or 17.

- 29. A part of a plant obtained from a plant of any one of the claims 22 to 25 wherein said part contains increased levels of trehalose.
- 35 30. A part according to any one of the claims 27 to 29 selected from the group consisting of bulbs, flowers, fruits, hairy roots, leaves, microtubers, pollen, roots, seeds,

- 51 -

stalks and tubers.

25

35

31. A method of preserving a plant or plant part in the presence of trehalose, comprising the steps of:

- 5 (1) growing a plant of any one of the claims 25 to 26, or growing a plant part of any one of the claims 29 to 30,
 - (2) harvesting the plant or the plant part which contains trehalose, and
 - (3) air drying the plant or plant part or alternatively,
- 10 (4) freeze drying the plant or plant part.
 - 32. A dried plant or plant part which obtainable by the method of claim 31.
- 15 33. A method for the production of trehalose comprising the steps of:
 - (1) growing a plant of claim 23 under conditions allowing for the production of trehalose,
 - (2) harvesting said plant or a part thereof,
- 20 (3) isolating the trehalose from the said plant or the said part thereof.
 - 34. Trehalose which is substantially free from bacterial or yeast contaminants.

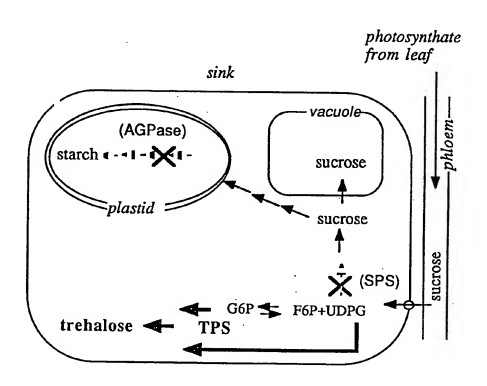
35. An isolated DNA sequence encoding a trehalose phosphate synthase activity.

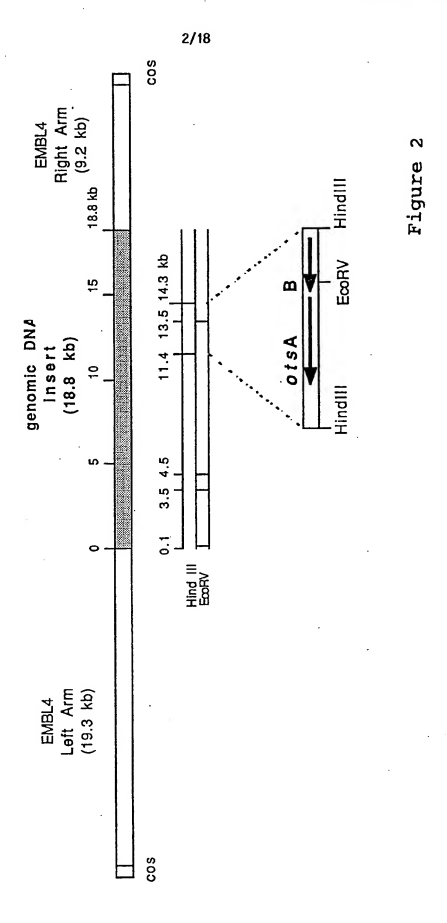
- 36. An isolated DNA sequence according to claim 34, 30 which is obtained from <u>E. coli</u>.
 - 37. An isolated DNA sequence according to claim 35 which is represented by SEQIDNO: 2, or an isolated DNA sequence hybridising therewith under stringent conditions.
 - 38. An isolated nucleic acid sequence that codes for the amino acid sequence of SEQIDNO: 3.

FIGURE 1.

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ENGINEERING OF TREHALOSE-PRODUCTION IN PLANTS







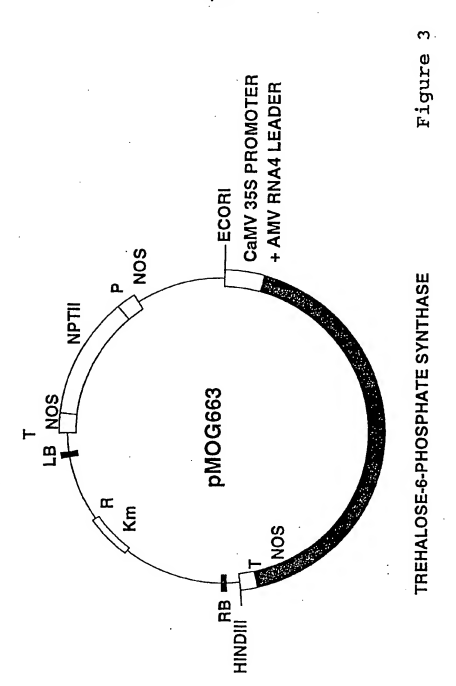
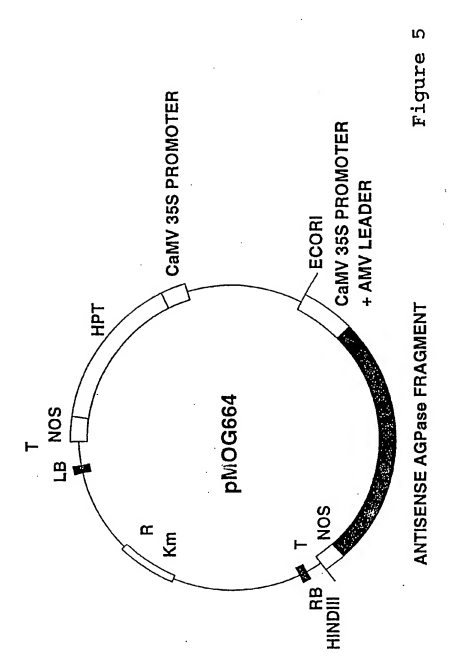
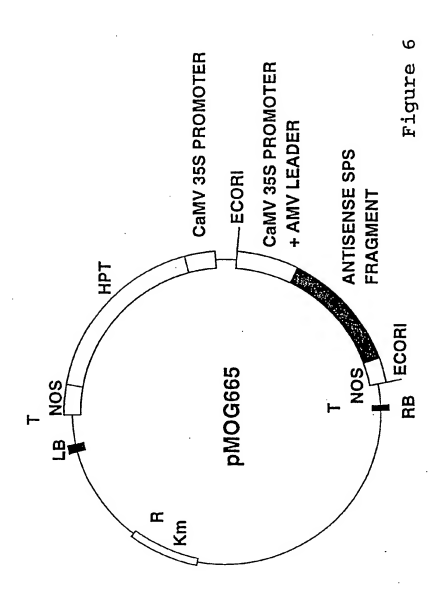
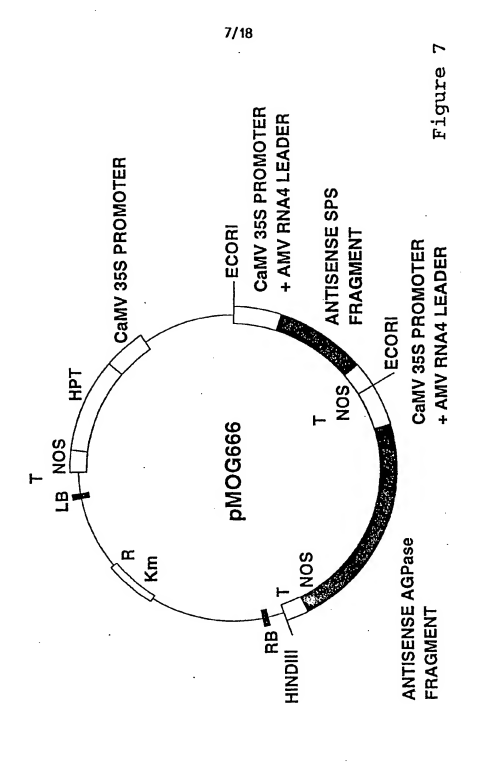


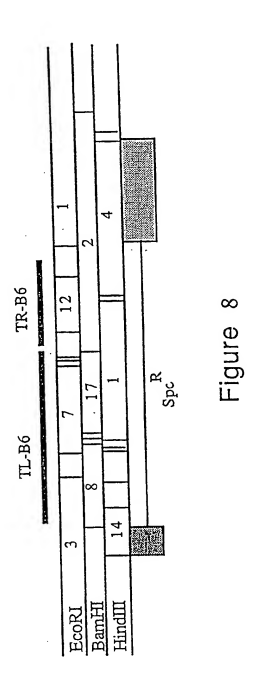
Figure 4

	09	120	180	240	300	360	370	
9	AGCA	CCCA	FTTG	AAGA	3GCT	CTAT		9
20 1 60	TGCTCG	CACATC	TGATGC	TTCCAA	GGCAAG	TGGGCA		09 05
50	AGCT	AGAT	CATC	BATA	TAI	PACA		20
_	TAGTAG	CICCCC	CATGCCC	GCGGACCAGG TGACAAGATA TTCCAAAAGA 240	TTAAGCCACA TTGTGAATAT GGCAAGGGCT 300	ACGTGA		<u>.</u>
1 40	TATG	LTGA	CICI	CAGG	CACA	CCLT		1 40
_	GTGAAG	51 CTTGCAAACA TGAAAGGAGT TCACCGAGTT GATCTCTTGA CTCGGCAGAT CACATCCCCA 120	21 GAGGITGAIT CTAGCIAIGG IGAGCCAAIT GAGAIGCICI CAIGCCCAIC IGAIGCIITG 180	GCGGAC	TTAAGC	GIGTGGCCIT ACGIGATACA IGGGCACTAT 360		_
30	CCAG	CTT	ATT	CCT	AGCA	AGCA	•	30
-	AGGTGG	TCACCG	TGAGCC7	TCGGAT	TGATGG!	TGGAAA	,	20 30
20	ATAC	GAGT	ATGG	CTAT	TTGT	ATGC		20
1 10 . 1 20	ATTCTG	TGAAAG	CIAGCI	TGCCTA	CAGAAT	AAGTCA		_
10	GTG	ACA	PATT	TGG	TAC	3AGC	CTG	10
_	CTAGGTC	CTTGCA	GAGGITC	31 GCTGCTGTGG TGCCTACTAT TCGGATCCCT	41 ATTTACATAC CAGAATTTGT TGATGGAGCA	11 ATAGGGGAGC AAGTCAATGC TGGAAAAGCA	51 GCCGATGCTG	1 10
	-1	51	21	31	러	7	51	









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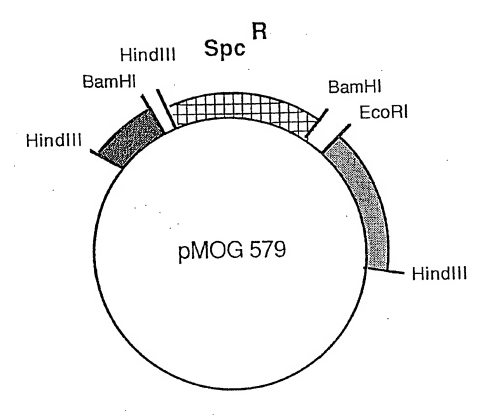


Figure 9

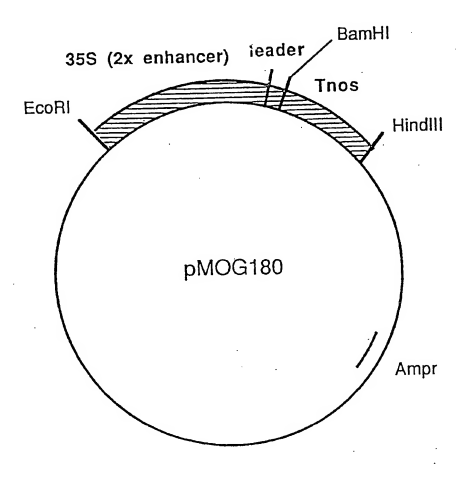


Figure 10

T /QT-									1	1								
H	GAG AAA ATA	ACA	GGA	ACA GGA GTG	ATG	ATG ACT ATG	ATG	ÀGT	CGI	CGT TIA GTC GIA	GTC	GTA	GTA	TCI	AAC	CGG	ATT	GC.№
					met	thr	met	ser	arg	arg leu	val	val val val		ser	asn	arg	ile	ala
									73/31	근								
	CCA CCA GAC	GAG	CAC	ပ္ပပ္ပ	GCC AGT	AGT	ပ္ပ	GCC GGT	ပ္ပင္ပ	CLL	ပ္ပပ္ပ	GTT	ပ္ပင္ပ	ATA	CIG	පුදුල	GCA	CIG
	asp	glu	his	ala	ala	ser	ala gly	gly	gly leu ala val gly ile leu	leu	ala	val	gly	ile		gly	ala	leu
	103/41					· ·			133/51	.21								
_	3CA	299	GGA	GGC GGA CTG TGG	TGG	TIT GGC TGG AGT GGT GAA ACA GGG AAT GAG	ပ္ပပ္ပ	TGG	AGT	GGT	GAA	ACA	999	AAT	GAG	GAT	CAG	ဗ္ဗ
	lys ala ala	gly	gly	leu	trp	phe	gly	trp	ser	ser gly glu thr	glu	thr	gly e	asn	glu asp	asb	gln	pro
									193/71	71								
	AAG	GTG	3	AAA	AAA GGT AAC ATT ACG TGG GCC TCT TTT AAC CTC	AAC	ATT	ACG	TGG	ပ္ပ	TCT	TII	AAC	CIC	AGC GAA	SAA	CAG	GAC
	lys	val	<u>ې</u>	lys c	gly	asn ile	ile	thr	trp ala	ala	ser	phe	asn	leu	ser	3]n	gln	asp
									253/91	,91								
_	CTT GAC GAA	TAC	TAC	TAC AAC CAA TTC TCC AAT GCC GTT CTC TGG	CAA	TTC	TCC	AAT	gu	GTT	CIC	\mathbf{TGG}	ပ္ပပ္ပ	GCT	$\mathtt{T}\mathtt{T}\mathtt{I}$	CAT	TAT	ეეე
leu asp c	glu	tyr	tyr	asn	gln	phe	ser	asn	asn ala val leu trp	val	leu	trp	pro	ala phe		his	tyr.	arg
									313/111	111								
CTC GAT (TE	GTG	CAA	GTG CAA TIT CAG CGT CCT GCC TGG GAC GGC TAT CTA CGC	CAG	CGT	CCI	ပ္ပပ္ပ	TGG	GAC	ggg	TAT	CIA	ပ္ပင္ပင	GTA AAT GCG	AAT	900	TIG
	leu	val	gln	phe	qln	arg	pro	ala	trp	asp	gly	tyr	leu	arg	val	asn	ala leu	leu

FIG. 11 A (Cont.)

	TAT	tyr		TIC	phe		ACC	thr		CTG	1en		CAT	his		GAA	glu		CTG	len
	GAT	asb		GGT	gly		GAC	asp		CGI	arg		AGC	ser		AAA	lys		GAA	glu
	CAC	his		ALT	ile		TAT	tyr		GAT	asp		AGC GCA AAA	1ys		ပ္ပင္ပင္ပ	pro		ဗ္ဗဗ္ဗ	ala
	ATC	ile		ပ္ပပ္ပ	arg		ACA	thr		AAC	asn		GCA	ala		GAA	glu		CAG GCT GCC GGG CCA CTG CCG CCA AAA CTG GCG CAA CTT AAA GCG	lys
	ŢĞĞ	asp asp asp ile ile trp		AAT	asn		900	pro		GAA	glu		AGC	ser		CCG ATC GGC ATT	gly ile		CTT	leu
	ATC	ile		AAT	asn		CIG	leu		ACA	thr		CGT	arg		ပ္ပ	gly		CAA	gln
	ATT	ile		GTG	val			ala		CAG	gln		ACG ACA	thr		ATC	ile		ggg	ala
	GAC	asp		GGA	gly		AAC GCG	asn		TIC	phe		ACG	thr			pro	•	CTG	leu
131	GAT	asp	151	AAA CGG	lys arg	171	ATC TTC	ile phe	191	CTG GGT	leu gly	211	CGC GTC	arg val	673/231	GTC TAC	tyr	733/251	AAA	lys
3/3/131	GAC	asp	433/151		lys	493/171	ATC	i: 1e	553/191	CIG	leu	613/211	ပ္ပဗ္ဗ		673/			733,	CCA	pro
	CAA	gln		ပ္ပင္ပ	arg		CCG GAA	glu		TTG	leu		ACC	thr		CGA ACA GAA	glu		SCG	pro
	TTG	leu		TTA	leu		ပ္သည္သ	pro		GAT TTG	asp		TCT AAC CTG ACC	leu		ACA	thr		CTG	leu
	AAA TTA CTG CCG CTG	leu		GAA	glu		TT CCT TTC CCG ACA	thr		TAT	tyr		AÀC	asn		CGA	arg		CCA	pro
	CCG	pro		CAT	his		SCG	pro		GAT	asp		TCI	ser		TTT	phe		999	414
	CTG	len		ပ္ပ	ala		TTC	phe		TGT	ςλs		CTT	leu		GCA	ala		ပ္ပ	ala
	TTA	leu		TII	apa		CCT	pro		CTT	len		TGI	cys		AAA	lys		GCT	ala
	AAA	lys		CCA			A I:	•~		CAG	gln		GAT	asp cys		ပ္ပ္ပ	gly		CAG	gln
	GAT	asp		TTG	leu		CAT	his		GAA	glu		CTG	leu		\mathbf{TGG}	trp		AAA	>
121	GCA	leu ala asp	141	CTG	leu	161	CIG	1eu	181	CTT	1eu	/201	TTC	phe	/221	CCC	ala	/241	ggg	, (0
343/121	CIG	leu	403/	CAC	his	463/	TTT	phe	523/	TIG	leu	583/	909	ala	643,	ACA	thr	703,	ATA	1.0

FIG. 11 B (Cont.)

	GAG	glu		CGI	arg		CAT	his		ပ္ပင္ပ	pro		TCT	ser		GTT	val		GCA	ala
	CCA	pro		ATT	ile		CGT	arg		ACG	thr		TAC	tyr		TAT	tyr		පුදුල	ala
	TIG	len		AAA	lys		ATT	ile		TGG	trp		ပ္ပပ္ပ	arg		GTA GCA AAA GAG	glu		GGA	gly
	GGT	g1y		GGT	gly		GAT	asp		ဥဌဌ	gly		TIC	phe		AAA	lys		gag	ala
	AAA	lys		CAT	his		CAG	gln		TTA	leu		AAA ATA	lys ile		GCA	ala		TTT	phe
	ICC	ser		CAT	his		TAT	tyr		CAA	gln		A.Y.			GIA	val		CAA	gln
	TAT	tyr		CAG	gln		GCC	ala		GGG	gly		ATG	met		GGG ATG AAC CTG	leu		TCG	ser
	GAT	asp		CCG	pro		CAA	gln		TAC	gly lys tyr		AAA TTA CTG	lys leu leu		AAC	gly met asn		CIT GIT CIT	leu
T / 7	CIG	leu	291	TAT	tyr	311	GAT GTG		331	GGT AAA	$_{ m lys}$	1033/351	TTA	leu	1093/371	ATG	met	1153/391	GTT	val
172/56/	CGG	arg	853/291	AAA	lys	913/311	GAT	asp val	973/331			1033	AAA	1ys	1093			1153	CTT	gly val leu val leu
	GAA	glu		GAA	glù		GGT	gly		AAT	asn		CGT	arg		GAC	asp		GTT	val
	GIC	val		CIG	leu		cgT	arg.		ATT	ile		GAC	asp		CGI	arg		ပ္ပ	gly
	TCT	ser		TIG	leu		TCG	ser		CGA	arg		TTT	phe		CIG	len		ညည	asn pro
	$_{ m LLL}$	phe		ggg	ala		ACG	thr		GGA	gly		CAT	his		CCA	thr pro		AC CCA GCC AAT	asn
	ATC	ile		GAA	glu		CCA	pro		GCI	ala		CAG	gln		ACG	thr		ည္ဟ	ala
	CAA AAT	asn		TAT	tyr		TT GCA	ala		GAA	glu		TG AAT	asn		GTG	val		CCA	asp pro
	CAA	gln		ည္ပ	ala		Æ	ile		AAT	asn		₽	leu		TTA	leu		O	asp
	GTA	val			leu		CAG	gln		GAA	glu		TAT	tyr		9	gly		CAG	gln
261	AAC	asn	281	TTT	phe	301	ACC	thr	321	CIC	leu	/341	TAT	tyr	/361	GTG	asp val gly	1123/381	GCT	ala
763/261	AAA AAC	lys	823/281	CGT	arg phe	883/	TAT	tyr thr gln	943/	CAG	gln	1003	CTT	leu	1063	GAC	asp	1123	GCT GCT CAG	ala

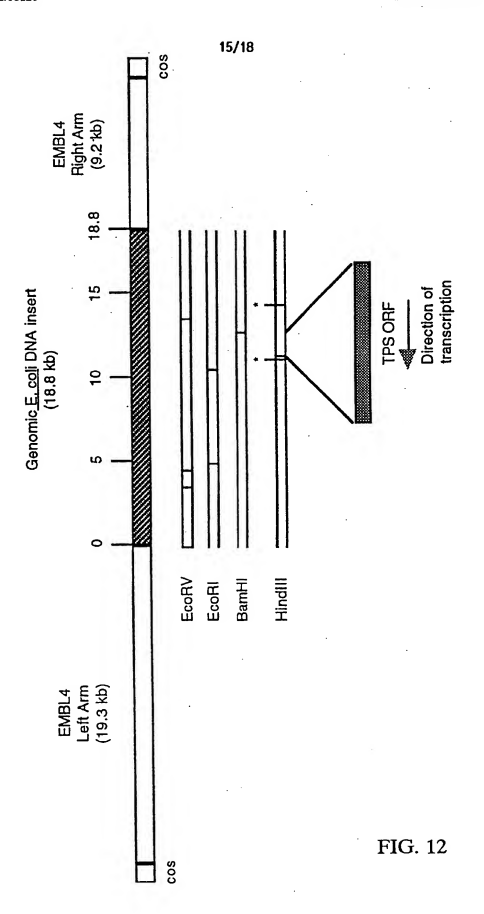
FIG. 11 C (Cont.)

CTGAAG leu trp gln glu cys phe ile ser asp leu lys CAG CGC GAT AAA GTT GCT ACC TTT CCA AAG pro ATT AGC GAC CTA GAA CGT ATT TCC CGT CAT GCA GAA ATG CCC TAC GAT CGT GAC GAA GTT GCA GCT glu val ala ala glu met phe his ala thr TTC pro tyr asp arg asp arg gln arg asp lys val glu arg ile ser TGG CAG GAG TGC 1273/431 1393/471 1333/451 asn ala glu ser,gln AAC GAG TTA ACG TCG GCG TTA ATT GTT AAC CTG GAT CGT GCA TTG ACT ATG TCG CTG GCG ala asp val ile val lys asn asp ile asn his CAG ATA GIT CCG CGA AGC GCG GAA AGC CAG GAC GIT AIC GIG AAA AAC GAI AIT AAC CAC len ala leu ile val ser thr met pro arg ser leu ser. asn glu leu thr leu asp arg ala gln ile val 1303/441 1243/421 1363/461 1423/481

1213/411

1183/401

FIG. 11 D



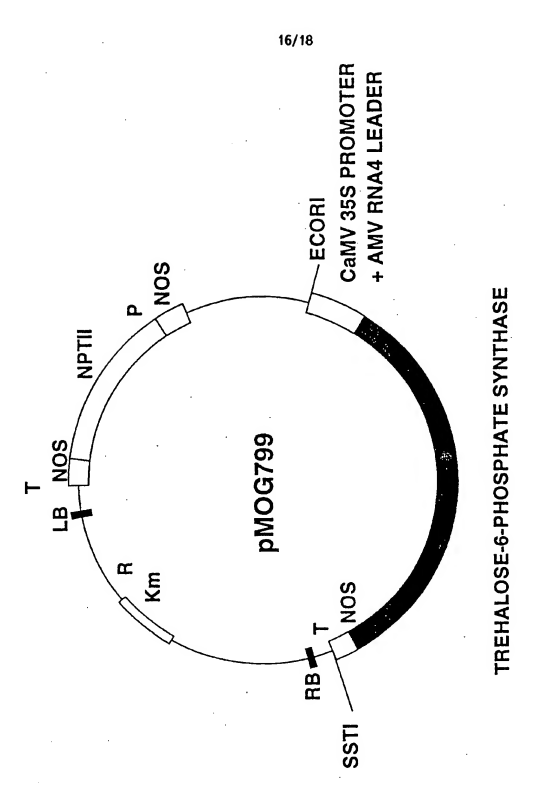


FIG. 13

PCT/EP93/02290

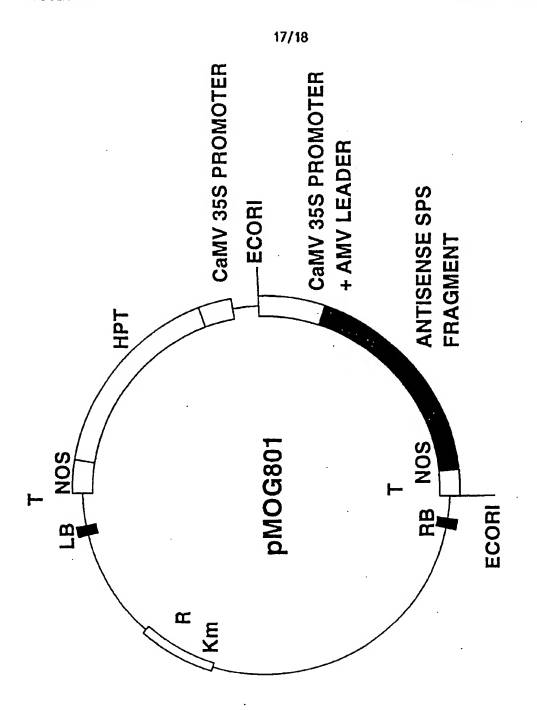


FIG. 14

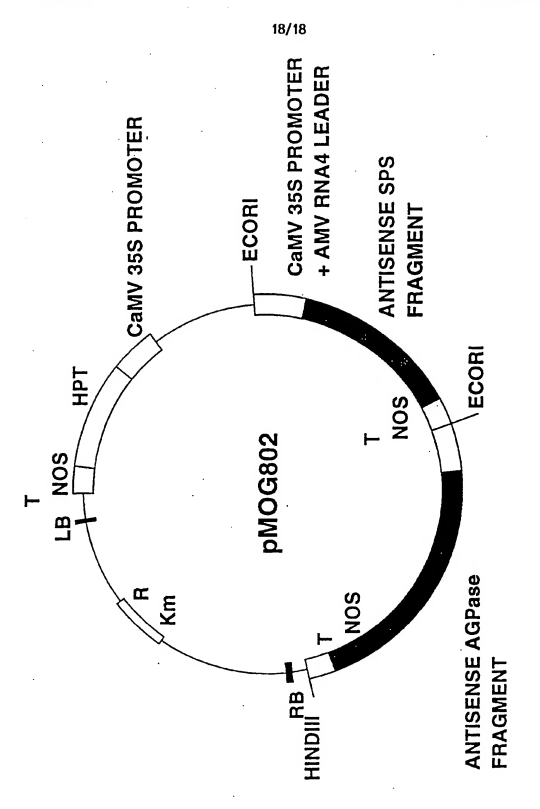


FIG. 15

Inter xonal Application No
PCT/EP 93/02290

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Documentat	tion searched other than m	inimum documentation	to the extent that such o	documents are included i	n the fields searched
Electronic d	ata base consulted during	the international search	(name of data base and	, where practical, search	terms used)
C. DOCUM	IENTS CONSIDERED T	O BE RELEVANT			
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	nation) DOCUMENTS CONSIDERED TO BE RELEVANT	
ategory *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
	EUROPEAN JOURNAL OF BIOCHEMISTRY vol. 209, no. 3, November 1992 pages 951 - 959 BELL, W., ET AL. 'Characterization of the 56-kDa subunit of yeast trehalose-6-phosphate synthase and cloning of its gene reveal its identity with the product of CIF1, a regulator of carbon catabolite inactivation' see the whole document	34
X	YEAST vol. 8 , 1992 pages 183 - 192 GONZALES, M.I., ET AL. 'Molecular cloning of CIF1, a yeast gene necessary for growth on glucose' see the whole document	34
X	J. BACTERIOLOGY vol. 174, no. 3 , February 1992 pages 889 - 898 KAASEN, I., ET AL. 'Molecular cloning and physical mapping of the otsBA gene, which encode the osmoregulatory trehalose pathway of Escherichia coli: Evidence that transcription is activated by KatF (AppR)' cited in the application see the whole document	34-38
X · _	EMBL SEQUENCE DATABASE REL. ACC. NO. X69160 27 May 1993	35-38
A	EP,A,O 451 896 (GIST-BROCADES) 16 October 1991 see the whole document	1-38
A	CURRENT BIOLOGY vol. 2, no. 11 , 1992 pages 594 - 596 TOMOS, D. 'Life without water' see page 596, left column, last paragraph	1-38
	PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA vol. 89 , April 1992 , WASHINGTON US pages 2600 - 2604 TARCZYNSKI, M.C., ET AL. 'Expression of a bacterial mtlD gene in transgenic tobacco leads to production and accumulation of mannitol' see the whole document	1-38

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		PCT/EP 9	3/02230
	ation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the relevant passages		Relevant to claim No.
A .	PLANT PHYSIOLOGY. vol. 99, no. 1 , May 1992 , ROCKVILLE, MD, USA. page 27 TARCZYNSKI, M.C., ET AL. 'Evaluation of the concept of osmoprotection : Effect of mannitol production in transgenic tobacco' see abstract 162		1-38
A	J. BIOTECHNOLOGY vol. 7, no. 1 , 1988 pages 23 - 32 COUTINHO, C., ET AL. 'Trehalose as cryoprotectant for preservation of yeast strains' see the whole document		31,32
E	WO,A,93 17093 (OY ALKO) 2 September 1993 see the whole document		34
E	EP,A,O 577 915 (ALGIST-BRUGGEMAN) 12 January 1994 see page 7, line 50 - line 54		34
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Inte: onal Application No
PCT/EP 93/02290

Patent document cited in search report	Publication date	Patent memb		Publication date
EP-A-0451896	16-10-91	AU-B- AU-A- JP-A-	636022 7378291 5184353	08-04-93 03-10-91 27-07-93
WO-A-9317093	02-09-93	AU-B-	3500993	13-09-93
EP-A-0577915	12-01-94	NONE		

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